

US010010507B1

## (12) United States Patent

Chong et al.

### (10) Patent No.: US 10,010,507 B1

(45) **Date of Patent: Jul. 3, 2018** 

## (54) PHARMACEUTICAL FORMULATIONS OF A BRUTON'S TYROSINE KINASE INHIBITOR

- (71) Applicant: **Pharmacyclics LLC**, Sunnyvale, CA
- (72) Inventors: Ching W. Chong, Fremont, CA (US); Robert Kuehl, San Francisco, CA
  - (US); **Heow Tan**, Cupertino, CA (US); **Harisha Atluri**, Palo Alto, CA (US)
- (73) Assignee: **Pharmacyclics LLC**, Sunnyvale, CA (US)
- (\*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.
- (21) Appl. No.: 15/909,779
- (22) Filed: Mar. 1, 2018

#### Related U.S. Application Data

- (63) Continuation of application No. 15/862,995, filed on Jan. 5, 2018, which is a continuation of application No. 15/467,414, filed on Mar. 23, 2017, now abandoned, which is a continuation of application No. 15/060,010, filed on Mar. 3, 2016, now Pat. No. 9,655,857.
- (60) Provisional application No. 62/193,518, filed on Jul. 16, 2015, provisional application No. 62/127,717, filed on Mar. 3, 2015.
- (51) Int. Cl.

  A61K 31/52 (2006.01)

  A61K 9/20 (2006.01)

  A61K 31/519 (2006.01)
- (52) U.S. CI.
  CPC ............ A61K 9/2054 (2013.01); A61K 9/2009
  (2013.01); A61K 9/2013 (2013.01); A61K
  9/2018 (2013.01); A61K 9/2027 (2013.01);
  A61K 9/2095 (2013.01); A61K 31/519

(2013.01)

#### (56) References Cited

#### U.S. PATENT DOCUMENTS

4,824,982	A	4/1989	Vahlensieck et al
5,033,252	A	7/1991	Carter
5,052,558	A	10/1991	Carter
5,145,684	A	9/1992	Liversidge et al.
5,323,907	A	6/1994	Kalvelage
5,397,787	A	3/1995	Buzzetti et al.
5,593,997	A	1/1997	Dow et al.
6,160,010	A	12/2000	Uckun et al.
6,221,900	B1	4/2001	Uckun et al.
6,303,652	B1	10/2001	Uckun et al.

6,306,897	B1	10/2001	Uckun et al.
6,326,469	B1	12/2001	Ullrich et al.
6,410,054	B1	6/2002	Thosar et al.
6,506,769	B2	1/2003	Snow et al.
6,660,744	B1	12/2003	Hirst et al.
6,753,348	B2	6/2004	Uckun et al.
6,770,639	B2	8/2004	Snow et al.
6,921,763	B2	7/2005	Hirst et al.
7,138,420	B2	11/2006	Bentzien et al.
7,332,497	B2	2/2008	Hirst et al.
7,514,444	B2	4/2009	Honigberg et al.
7,718,662	B1	5/2010	Chen et al.
7,732,454	B2	6/2010	Verner
7,741,330	В1	6/2010	Chen et al.
7,825,118	B2	11/2010	Honigberg et al.
7,960,396	B2	6/2011	Honigberg et al.
8,008,309	B2	8/2011	Honigberg et al.
8,088,781	B2	1/2012	Honigberg et al.
8,124,126	B2	2/2012	Bosse et al.
8,158,786	B2	4/2012	Honigberg et al.
8,232,280	B2	7/2012	Honigberg et al.
8,236,812	B2	8/2012	Honigberg et al.
8,306,897	B2	11/2012	Yolles
8,377,946	B1	2/2013	Chen et al.
8,399,470	B2	3/2013	Honigberg et al.
8,399,471	B2	3/2013	Figueroa Perez et al
8,476,284	B2	7/2013	Honigberg et al.
8,497,277	B2	7/2013	Honigberg et al.
8,501,724	B1	8/2013	Chen et al.
8,501,751	B2	8/2013	Honigberg et al.
8,552,010	B2	10/2013	Honigberg et al.
8,557,803	B2	10/2013	Yamamoto et al.
8,563,563	B2	10/2013	Honigberg et al.
8,633,311	B2	1/2014	Bommer et al.
8,642,604	B2	2/2014	Knight et al.
8,658,653	B2	2/2014	Honigberg et al.
8,691,546	B2	4/2014	Honigberg et al.
		(Con	tinued)
		( - 011	<i>-</i> )

#### FOREIGN PATENT DOCUMENTS

CN 101610676 A 12/2009 CN 103121999 A 5/2013 (Continued)

#### OTHER PUBLICATIONS

ACS 2015 (http://www.cancer.org/cancer/non-hodgkinlymphoma/detailedguide/non-hodgkin-lymphoma-types-of-non-hodgkinlymphoma).

(Continued)

Primary Examiner — Raymond J Henley, III (74) Attorney, Agent, or Firm — Foley Hoag LLP

#### (57) ABSTRACT

Described herein are pharmaceutical formulations of Bruton's tyrosine kinase (Btk) inhibitor 1-((R)-3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl)piperidin-1-yl)prop-2-en-1-one. Also disclosed are methods of using the Btk inhibitor, alone or in combination with other therapeutic agents, for the treatment of autoimmune diseases or conditions, heteroimmune diseases or conditions, cancer, including lymphoma, and inflammatory diseases or conditions.

#### 28 Claims, 3 Drawing Sheets

# US 10,010,507 B1 Page 2

(56)	Referer	ices Cited	2010/0022561	A1	1/2010	Honigberg et al.
II:	S PATENT	DOCUMENTS	2010/0041677 2010/0254905			Honigberg et al. Honigberg et al.
0.	O. 171111111	DOCOMENTS	2010/0324050			Honigberg et al.
8,697,711 B2	2 4/2014	Honigberg et al.	2011/0039868			Honigberg et al.
8,703,780 B2		Honigberg et al.	2011/0086866 2011/0160273			Chen et al. Buschmann et al.
8,735,403 B2 8,735,404 B2		Honigberg et al. Honigberg et al.	2011/0177011			Currie et al.
8,741,908 B2		Honigberg et al.	2011/0224235			Honigberg et al.
8,748,438 B2		Honigberg et al.	2011/0281322 2012/0065201			Honigberg et al. Honigberg et al.
8,748,439 B2 8,754,090 B2		Honigberg et al. Buggy et al.	2012/0003201			Buggy et al.
8,754,090 B2		Honigberg et al.	2012/0087915	A1	4/2012	Buggy et al.
8,759,516 B2	2 6/2014	Honigberg et al.	2012/0088912			Honigberg et al.
8,809,273 B2		Honigberg et al.	2012/0095026 2012/0100138			Honigberg et al. Buggy et al.
8,883,435 B2 8,883,803 B2		Honigberg et al. Honigberg et al.	2012/0101113		4/2012	Honigberg et al.
8,940,750 B2	2 1/2015	Honigberg et al.	2012/0101114			Honigberg et al.
8,952,015 B2		Honigberg et al.	2012/0108547 2012/0108612			Jankowski et al. Honigberg et al.
8,957,079 B2 8,975,266 B2		Honigberg et al. Honigberg et al.	2012/0115889			Honigberg et al.
8,987,233 B2		Pan et al.	2012/0122894		5/2012	Honigberg et al.
8,999,999 B2		Buggy et al.	2012/0129821 2012/0129873			Honigberg et al. Honigberg et al.
9,012,463 B2 9,079,908 B2		Chen et al. Honigberg et al.	2012/0125873			Honigberg et al.
9,107,924 B2		Buggy et al.	2012/0165328	A1		Honigberg et al.
9,117,924 B2	2 8/2015	Kitagawa et al.	2012/0178753		7/2012	Honigberg et al.
9,125,889 B2		Buggy et al.	2012/0183535 2012/0184013			Buggy et al. Honigberg et al.
9,127,012 B2 9,127,069 B1		Honigberg et al. Tabuteau	2012/0184567			Honigberg et al.
9,133,198 B2		Honigberg et al.	2012/0202264			Honigberg et al.
9,133,201 B2		Honigberg et al.	2012/0214826 2012/0238733			Honigberg et al. Vasudevan et al.
9,133,202 B2 9,139,591 B2		Honigberg et al. Honigberg et al.	2012/0252821			Honigberg et al.
9,181,257 B2		Honigberg et al.	2012/0252822		10/2012	Honigberg et al.
9,181,263 B2	2 11/2015	Honigberg et al.	2012/0277177			Amparo et al.
9,193,735 B2	2 11/2015	Honigberg et al. Honigberg et al.	2012/0277225 2012/0277254			Honigberg et al. Honigberg et al.
9,206,189 B2 9,212,185 B2		Honigberg et al.	2012/0277255		11/2012	Honigberg et al.
9,266,893 B2		Honigberg et al.	2012/0283276			Honigberg et al.
9,273,051 B2		Chen et al.	2012/0283277 2012/0296089			Honigberg et al. Honigberg et al.
9,278,100 B2 9,296,753 B2		Honigberg et al. Smyth et al.	2012/0328679			Curatolo et al.
9,409,911 B2		Honigberg et al.	2012/0329130			Honigberg et al.
9,540,382 B2		Purro et al.	2013/0005745 2013/0005746			Honigberg et al. Honigberg et al.
9,545,407 B2 9,655,857 B2		Shu et al. Chong et al.	2013/0012525			Honigberg et al.
9,713,617 B2		Purro et al.	2013/0018060		1/2013	Honigberg et al.
9,725,455 B		Purro et al.	2013/0035334 2013/0079327			Honigberg et al. Yamamoto et al.
9,828,383 B1 2002/0016460 A		Purro et al. Snow et al.	2013/0079327			Chen et al.
2002/0010400 A		Wells et al.	2013/0195852		8/2013	Buggy et al.
2003/0013125 A		Braisted et al.	2013/0202611		8/2013	Buggy et al.
2003/0035833 A			2013/0273030 2013/0310402		11/2013	Buggy et al. Buggy et al.
2003/0040461 A 2003/0118078 A		McAtee Carlson et al.	2013/0338172		12/2013	Smyth et al.
2003/0125235 A	1 7/2003	Foxwell	2014/0039186			Honigberg et al.
2003/0225144 A		Woodward et al.	2014/0057907 2014/0079690			Honigberg et al. Buggy et al.
2004/0006083 A 2005/0008640 A		Hirst et al. Waegell et al.	2014/0080844			Chen et al.
2005/0084905 A		Prescott et al.	2014/0088116			Morales Rojas et al.
2005/0090499 A		Currie et al.	2014/0128413 2014/0128414			Honigberg et al. Honigberg et al.
2005/0101604 A 2005/0153990 A		Currie et al. Watkins	2014/0135347			Honigberg et al.
2005/0196851 A		Uckun	2014/0142123	A1	5/2014	Honigberg et al.
2005/0209255 A		Jimenez et al.	2014/0163027			Verner et al.
2006/0079494 A		Santi et al. Uckun et al.	2014/0163046 2014/0171453			Honigberg et al. Honigberg et al.
2006/0167090 A 2007/0065449 A		Verschraegen	2014/0187564			Honigberg et al.
2007/0105136 A	1 5/2007	Staudt et al.	2014/0187565			Honigberg et al.
2007/0281907 A		Watkins	2014/0194446			Buggy et al.
2008/0076921 A 2008/0108636 A		Honigberg et al. Honigberg et al.	2014/0212485 2014/0243355			Honigberg et al. Honigberg et al.
2008/0214501 A		Pan et al.	2014/0288037			Casebier et al.
2008/0242707 A	1 10/2008	Schuckler et al.	2014/0336203	A1	11/2014	Smyth et al.
2009/0010911 A		Andreotti et al.	2014/0336206			Honigberg et al.
2009/0105209 A 2009/0186898 A		Dewdney et al. Dewdney et al.	2014/0377258 2014/0378446			Stern et al. Chen et al.
2009/0180898 A 2009/0317836 A		Kuhn et al.	2015/0018336			Chen et al.
2005.0517050 11.	- 12/2009				2.2010	

U.S. PATENT DOCUMENTS	(56)	Refere	nces Cited	WO	WO-2005/037843		4/2005	
2015/0031710   Al   1.2015   Baggy et al.   WO   WO-2006/003577   Al   2.2006		II C DATENE	E DOCUMENTS	WO			7/2005	
2015/03/1710   Al   12/015   Baggy of al   WO   WO-2006/03/274   Al   20/06		U.S. PATENT	DOCUMENTS					
2015 0015711 Al 1   12015   Boggy cal.	2015	/0021710 A1 1/2015	Puggy et al					
2015 003818 Al   22015   Balsaubarmanian				WO				
2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015		0038518 A1 2/2015	Balasubramanian					
2015 033661   A1   \$2015   Honjeberg et al.   WO   WO-2007 003225   A1   12007   2015 032161   A1   62015   Puro et al.   WO   WO-2007 003225   A2   12007   2015 032160   A1   \$2015   Wong   WO   WO-2007 003225   A2   52007   2015 032160   A1   \$2015   Susas et al.   WO   WO-2008 003218   A2   42008   2015 032107   A1   \$2015   Egusa et al.   WO   WO-2008 003218   A2   42008   2015 0325 037   A1   \$2015   Egusa et al.   WO   WO-2008 003218   A2   42008   2015 0326361   A1   \$2015   Byrd et al.   WO   WO-2008 1974   A2   12009   2015 0326361   A1   \$2015   Byrd et al.   WO   WO-2008 1374   A2   12009   2015 0326361   A1   102015   Honjeberg et al.   WO   WO-2008 1374   A2   12009   2015 0326361   A1   102015   Honjeberg et al.   WO   WO-2009 13237   A1   12209   2016 000773   A1   22015   More et al.   WO   WO-2009 13237   A1   12209   2016 000773   A1   22016   Bugy et al.   WO   WO-2009 13237   A1   12209   2016 000777   A1   22016   Bugy et al.   WO   WO-2009 13237   A1   12209   2016 000777   A1   22016   Bugy et al.   WO   WO-2009 13237   A1   12209   2016 000777   A1   22016   Bugy et al.   WO   WO-2009 13237   A1   12209   2016 000777   A1   22016   Bugy et al.   WO   WO-2009 13237   A1   12209   2016 000777   A1   22016   Bugy et al.   WO   WO-2010009342   A3   12010   2016 000777   A1   22017   Chen et al.   WO   WO-201000788   A2   12010   2017 0002090   A1   12017   Chen et al.   WO   WO-2010040780   A1   122011   2017 0007891   A1   22017   Chen et al.   WO   WO-2010040780   A1   122011   2017 0007891   A1   22017   Chen et al.   WO   WO-2010040780   A1   122011   2017 0007891   A1   22017   Chen et al.   WO   WO-2010040780   A1   122011   2017 0007891   A1   22017   Chen et al.   WO   WO-2010040780   A1   122011   2017 0007891   A1   22017   Chen et al.   WO   WO-2010040780   A1   122011   2017 0007891   A1   22017   Chen et al.   WO   WO-2010040780   A1   122011   2017 0007891   A1   22017   Chen et al.   WO   WO-2010040780   A1   122011   2017 0007891   A1   22017   Chen et al.   WO   WO-2010040		/0110871 A1 4/2015	Wong					
Display   Disp			Levy et al.					
2015 035871 Al   c.2015   Purro et al								
2015/02/24069   Al   2015   Egusset al   WO   WO-2007/18/790 & 12/0007   Egusset al   WO   WO-2007/18/790 & 12/0007   WO-2007/1								
2015/023407 A1   x2015   guss at al.								
2015/02/03/03/03   3   2015   1   1   2015   1   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015   2015	2015/							
2015/02/5618 Al 9/2015   Bonighery et al.   WO   WO-2008/108636 Al   9/2008   System of the composition of								
2015/02/75/1 Al 9-2015   Byrd et al.   WO   WO-2008/12/142 A2   10/2008								
2015/03/6752 Al 9/2015   Navarro								
2015/030610 Al   10/2015   Honipberg et al.   WO   WO-2099/151822 Al   4/2009								
2015/03/05/00 Al   02015   Honipberg et al.   WO   WO-2009/15/25/31 Al   12/2019   2016/000/05/31 Al   12/2015   Buggy et al.   WO   WO-2009/15/35/31 Al   12/2009   2016/000/37/31 Al   12/2016   Parket et al.   WO   WO-2010/05/38/32 Al   12/0016   WO   WO-2010/05/38/32 Al   12/0016   WO   WO-2010/05/38/34 Al   12/0016   WO   WO-2010/05/38/34 Al   WO   WO-2010/05/38/34 A								
2016/0006724   1   1/2016   Buggy et al.		/0306106 A1 10/2015	Honigberg et al.					
2016/06/274 Al   2016   Parket et al.								
2016/02/333 Al								
2016/02/3033 Al   \$2.016   Buggy et al.								
2016/025739 A   1								
2017/0002009 Al	2016/							
2017/0079981 Al 3-2017   Alluri et al.   WO WO-2011/152513 Al 12/2011   2017/00258729 Al 9/2017   Cohen et al.   WO WO-2011/162515 A2 12/2011   2017/00258729 Al 9/2017   Purro et al.   WO WO-2011/162515 A2 12/2011   2018/0072739 Al 3/2018   Goldman et al.   WO WO-2012/158764 Al 11/2012   2018/0072739 Al 3/2018   Goldman et al.   WO WO-2013/184572 Al 12/2013   2018/0072739 Al 3/2018   Goldman et al.   WO WO-2013/184572 Al 12/2013   2018/0072739 Al 3/2018   Goldman et al.   WO WO-2013/184572 Al 12/2013   2018/0072739 Al 3/2018   Goldman et al.   WO WO-2013/184572 Al 12/2013   2018/0072739 Al 3/2018   Goldman et al.   WO WO-2013/184572 Al 12/2013   2018/0072739 Al 3/2018   Goldman et al.   WO WO-2014/004707 Al 1/2014   2018/0072739 Al 3/2018   WO WO-2015/034478 Al 3/2015   2018/0072739 Al 3/2018   WO WO-2015/034478 Al 3/2015   2018/0072739 Al 3/2018   WO WO-2015/034878 Al 3/2015   2018/0072739 Al 1/2014   WO WO-2015/034878 Al 3/2015   2018/0072739 Al 1/2015   WO WO-2015/034878 Al 3/2015   2018/0072739 Al 1/2015   WO WO-2015/034878 Al 3/2015   2018/0072739 Al 1/2015   WO WO-2015/0314884 A 3/2015   2018/0072739 Al 1/2015   WO WO-2015/031488 Al 3/2015   2018/0072739 Al 1/2015   WO WO-2015/031488 Al 3/2015   2018/0072739 Al 1/2015   WO WO-2015/031488 Al 3/2015   2018/0072739 Al 1/2015   WO WO-2015/031888 Al 16/2015   2018/0072739 Al 1/2015   WO WO-2015/031888 Al 16/2015   2018/0072739 Al 1/2016   WO WO-2015/031888 Al 16/2015   2018/0072739 Al 1/2016   WO WO-2016/025729 Al 1/2016   2018/0072739 Al 1/2004   WO WO-2016/025729 Al 1/2016   2018/0072739 Al 1/2004   WO WO-2016/025042 Al 1/2016   2018/0072739 Al 1/2004   WO WO-2016/02504 Al 1/2016   2018/0072739 Al 1/2004   WO WO-2016/02504 Al 1/2016   2018/0072739 Al 1/2004   WO WO-2016/02504 Al 1/2016   2018/0072739 Al 1/2004   WO WO-2016/05042 Al 1/2016								
2017/0226/114 Al 8/2017   Cohen et al.								
2017/02/S8739   A1   9/2017   Purro et al.   WO   WO-2011/162515   A2   12/2011								
2017/03/05/19   Al   10/2017   Purro et al.   WO   WO-2012/15/8764   Al   11/2012								
2018/00/2733					WO-2012/021444	A1		
POREIGN PATENT DOCUMENTS		0028537 A1 2/2018	Atluri et al.					
FOREIGN PATENT DOCUMENTS								
FOREIGN PATENT DOCUMENTS	2018/	0072739 A1 3/2018	Goldman et al.					
FOREIGN PATERT   DOC UMENTS   WO   WO-2014/018567 A1   1/2014		EODEIGNI DATE	ENTE DOCLD TENTE					
CN         109323084 A 7/2014         WO WO-2015/038887 AI 3/2015           CN         104523695 A 4/2015         WO WO-2015/038887 AI 5/2015           CN         10508529 A 11/2015         WO WO-2015081180 AI 6/2015           CN         105294696 A 2/2016         WO WO-2015095772 A2 6/2015           CN         105440040 A 3/2016         WO WO-2015/1820110 A2 8/2015           CN         105640961 A 6/2016         WO WO-2015/145415 A2 10/2015           CN         105646484 A 6/2016         WO WO-2016/025720 AI 2/2016           CN         105646498 A 6/2016         WO WO-2016/025720 AI 2/2016           CN         105646499 A 6/2016         WO WO-2016/025720 AI 2/2016           CN         106619643 A 5/2017         WO WO-2016/0625720 AI 2/2016           CN         106619643 A 5/2017         WO WO-2016/050422 AI 2/2016           CN         106619643 A 5/2017         WO WO-2016/050422 AI 2/2016           CN         106619643 A 7/1989         WO WO-2016/050422 AI 2/2016           MO         WO-1997/028161 AI 3/1989         WO WO-2016/050422 AI 2/2016           WO WO-1997/049706 AI 2/2016         WO WO-2016/159588 AI 9/2016           WO WO-1997/049706 AI 2/2017         WO WO-2016/160598 AI 10/2016           WO WO-2009/040828 A2 10/1999         WO WO-2016/160598 AI 10/2016           WO WO-		FOREIGN PAIE	ENT DOCUMENTS					
CN	CN	103023084 A	7/2014					
CN								
CN         105294696 A 2 2/2016         WO WO-2015095772 A2 6/2015         6/2016           CN         105440961 A 6/2016         WO WO-2015/120110 A2 8/2015         8/2015           CN         105640848 A 6/2016         WO WO-2016/022942 A1 2/2016           CN         105646498 A 6/2016         WO WO-2016/0225720 A1 2/2016           CN         105646499 A 6/2016         WO WO-2016/025720 A1 2/2016           CN         105646499 A 6/2016         WO WO-2016/025720 A1 2/2016           CN         105646499 A 6/2016         WO WO-2016/025720 A1 2/2016           EP         1473039 A1 11/2004         WO WO-2016/05065 A1 2/2016           BP         H01167840 A 71/989         WO WO-2016/050422 A1 4/2016           BP         H01167840 A 71/989         WO WO-2016/127960 A1 8/2016           WO WO-1997/049706 A1 12/1997         WO WO-2016/127960 A1 8/2016           WO WO-1997/049706 A1 12/1997         WO WO-2016/1605088 A1 10/2016           WO WO-2006/000823         1/2000           WO WO-2006/000823         1/2000           WO WO-2001/01829 A2 3/2001         WO WO-2016/160604 A1 10/2016           WO WO-2001/041754         6/2001           WO WO-2001/041754         6/2001           WO WO-2002/08978 A2 5/2002         Advani et al. Effect of Btk inhibitor PCI-32765 is highly active and more proceedings (	CN							
CN								
CN								
CN								
CN 105646499 A 6/2016 CN 106619643 A 5/2017 EP 1473039 A1 11/2004 JP H01167840 A 7/1989 WO WO-20160050697 A1 2/2016 WO WO-2016/050422 A1 4/2016 JP H01167840 A 7/1989 WO WO-201679216 A1 5/2016 WO WO-1997/028161 A1 8/1997 WO WO-2016127960 A1 8/2016 WO WO-1997/049706 A1 12/1997 WO WO-2016139588 A1 9/2016 WO WO-1998/041525 A1 9/1998 WO WO-1998/041525 A1 9/1998 WO WO-2016/160598 A1 10/2016 WO WO-2000/000823 1/2000 WO WO-2001/055737 WO WO-2001/055737 WO WO-2001/019829 A2 3/2001 WO WO-2001/019829 A2 3/2001 WO WO-2001/041254 A1 6/2001 WO WO-2001/041258 A1 6/2001 WO WO-2001/044258 A1 6/2001 WO WO-2002/088797 A2 5/2002 WO WO-2003/001874 WO WO-2003/001854 WO WO-2004/096253 A1 11/2003 WO WO-2004/096258 A1 11/2004 WO WO-2004/096258 A1 11/2003 WO WO-2004/096258 A1 11/2004 WO WO-2004/09686 A2 11/2004 WO WO-2004/096258 A1 11/2004 WO WO-2004/096258 A1 11/2004 WO WO-2004/096258 A1 11/2004 WO WO-2004/096259 A1 11/2004 WO WO-2004/09688 A3 11/2004 WO WO-2005/005499 A1 1/2005								
CN 106619643 A 5/2017 EP 147303 A1 11/2004 WO WO-2016/050422 A1 4/2016 JP H01167840 A 7/1989 WO WO-2016/050422 A1 4/2016 JP 5841998 B2 1/2016 WO WO-1997/028161 A1 8/1997 WO WO-2016127960 A1 8/2016 WO WO-1997/049706 A1 12/1997 WO WO-2016139588 A1 9/2016 WO WO-1998/041525 A1 9/1998 WO WO-2016/160998 A1 10/2016 WO WO-1999/054286 A2 10/1999 WO WO-2000/006823 WO WO-2000/056737 WO WO-2001/056737 WO WO-2001/019829 A2 3/2001 WO WO-2001/019829 A2 3/2001 WO WO-2001/019829 A3 3/2001 WO WO-2001/041754 WO WO-2001/08829 A3 3/2001 WO WO-2001/08829 A3 6/2001 WO WO-2001/08829 A3 6/2001 WO WO-2001/08829 A3 6/2001 WO WO-2001/08829 A3 6/2001 WO WO-2001/08829 A3 16/2002 WO WO-2001/041754 WO WO-2001/088797 A2 5/2002 WO WO-2002/08797 A2 5/2002 WO WO-2002/089926 WO WO-2002/080926 WO WO-2003/008096 WO WO-2003/008096 WO WO-2003/046200 WO WO-2003/046200 WO WO-2003/046200 WO WO-2003/046200 WO WO-2003/0499A1 1/2003 WO WO-2003/0499A1 1/2003 WO WO-2004/096253 A1 11/2004 WO WO-2004/096253 A1 11/2004 WO WO-2004/0868 A2 11/2004 WO WO-2004/0868 A2 11/2004 WO WO-2004/0868 A3 11/2004 WO WO-2004/0868 A3 11/2004 WO WO-2005/00197 A2 1/2005 WO WO-2005/00197 A2 1/2005 WO WO-2005/00197 A2 1/2005 WO WO-2005/00199 A1 2/2005 WO WO-2005/00199 A1 2/2005 WO WO-2005/00199 A1 2/2005 WO WO-2005/00199 A1 1/2005 WO WO-2005/014599 A1 1/2005								
P								
F								
WO         WO-1997/028161         A1         8/1997         WO         WO-2016/139588         A1         9/2016           WO         WO-1997/049706         A1         12/1997         WO         WO-2016/160598         A1         10/2016           WO         WO-1999/054286         A2         10/1999         WO         WO-2016/160604         A1         10/2016           WO         WO-2000/000823         1/2000         WO         WO-2016/156127         A1         10/2016           WO         WO-2001/05238         10/2000         WO         WO-2001/019829         A3         3/2001           WO         WO-2001/019829         A3         3/2001         Advani et al. Effect of Btk inhibitor PCI-32765 monotherapy on responses in patients with relapsed aggressive NHL: Evidence of wO-2001/0441754         6/2001         responses in patients with relapsed aggressive NHL: Evidence of antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/038797         A2         5/2002         antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/076986         A1         10/2002         Supp):8012 (2010).           WO         WO-2003/060187         1/2003         Advani et al., "The Btk inhibitor PCI-32765 is highly active and well tolerated in patients (PTS) with relapsed/refractory B ce								
WO         WO-1997/049706         Al         12/1997         WO         WO-2016/160598         Al         10/2016           WO         WO-1998/041525         Al         9/1998         WO         WO-2016/160604         Al         10/2016           WO         WO-2000/00823         1/2000         WO         WO-2016/160604         Al         10/2016           WO         WO-2000/056737         9/2000         WO         WO-2001/019829         A3         3/2001           WO         WO-2001/019829         A3         3/2001         Advani et al. Effect of Btk inhibitor PCI-32765 monotherapy on responses in patients with relapsed aggressive NHL: Evidence of end wowled wo								
WO         WO-1998/041525 A1         9/1998         WO WO-2016/160604 A1         10/2016           WO         WO-1999/054286 A2         10/1999         WO WO-2016/160604 A1         10/2016           WO         WO-2000/000823         1/2000         WO-2016/1656127 A1         10/2016           WO         WO-2001/056737         9/2000         OTHER PUBLICATIONS           WO         WO-2001/019829 A2         3/2001         Advani et al. Effect of Btk inhibitor PCI-32765 monotherapy on responses in patients with relapsed aggressive NHL: Evidence of wO-2001/044258 A1         6/2001         responses in patients with relapsed aggressive NHL: Evidence of antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2001/044258 A1         6/2001         Annual Meeting Proceedings (Post-Meeting Edition), 28(15           WO         WO-2002/038797 A2         5/2002         Annual Meeting Proceedings (Post-Meeting Edition), 28(15           WO         WO-2002/076986 A1         10/2002         Advani et al., "The Btk inhibitor PCI-32765 is highly active and well tolerated in patients (PTS) with relapsed/refractory B cell           WO         WO-2003/040208         6/2003         malignancies: final results from a phase I study," Ann Oncol, 22(Supp 4):abstract 153 (2011).           WO         WO-2004/074290 A1         9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor Ibrutinib (PCI-32765) ha significant activi		WO-1997/049706 A1	12/1997					
WO         WO-2000/00823         1/2000         WO-2000/056737         9/2000           WO         WO-2001/025238         10/2000         OTHER PUBLICATIONS           WO         WO-2001/019829         A2         3/2001           WO         WO-2001/041754         6/2001         responses in patients with relapsed aggressive NHL: Evidence of antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/038797         A2         5/2002         Annual Meeting Proceedings (Post-Meeting Edition), 28(15)           WO         WO-2002/076986         A1         10/2002         Supp):8012 (2010).           WO         WO-2003/00187         1/2003         Advani et al., "The Btk inhibitor PCI-32765 is highly active and well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients (PTS) with relapsed/refractory B cell malignancies: final results from a phase I study," Ann Oncol, 22(Supp 4):abstract 153 (2011).           WO         WO-2003/097645         11/2003         22(Supp 4):abstract 153 (2011).           WO         WO-2004/096253         A1         11/2004           WO         WO-2004/100868         A2         11/2004           WO         WO-2005/000197         A2         1/2005           WO         WO-2005/005429         A1         1/2005           WO         WO-2005/00544								
WO         WO-2000/056737         9/2000           WO         WO-2001/025238         10/2000         OTHER PUBLICATIONS           WO         WO-2001/019829 A2         3/2001           WO         WO-2001/041754         6/2001         Advani et al. Effect of Btk inhibitor PCI-32765 monotherapy on responses in patients with relapsed aggressive NHL: Evidence of antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/038797 A2         5/2002         antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/38797 A2         5/2002         Annual Meeting Proceedings (Post-Meeting Edition), 28(15)           WO         WO-2002/076986 A1         10/2002         Supp):8012 (2010).           WO         WO-2003/090187         1/2003         Advani et al., "The Btk inhibitor PCI-32765 is highly active and well tolerated in patients (PTS) with relapsed/refractory B cell malignancies: final results from a phase I study," Ann Oncol, wowwww.com/support of the proceedings (POST) with relapsed/refractory B cell malignancies: final results from a phase I study," Ann Oncol, 22(Supp 4):abstract 153 (2011).           WO         WO-2004/096253 A1         11/2003         22(Supp 4):abstract 153 (2011).           WO         WO-2004/100868 A2         11/2004         Invitable PCI-32765 has significant activity in patients with relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.         Agathocleous et al.				WO	WO-2016156127	A1	10/2016	
WO         WO-2001/019829         A2 3/2001           WO         WO-2001/019829 A3 3/2001         Advani et al. Effect of Btk inhibitor PCI-32765 monotherapy on WO-2001/041754 6/2001           WO         WO-2001/041754 6/2001         fo/2001           WO         WO-2001/044258 A1 6/2001         responses in patients with relapsed aggressive NHL: Evidence of antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/38797 A2 5/2002         Annual Meeting Proceedings (Post-Meeting Edition), 28(15)           WO         WO-2002/076986 A1 10/2002         Supp):8012 (2010).           WO         WO-2003/080926 WO-2003/000187 1/2003         1/2003 Well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients from a phase I study," Ann Oncol, 22(Supp 4):abstract 153 (2011).           WO         WO-2003/046200 6/2003         malignancies: final results from a phase I study," Ann Oncol, 22(Supp 4):abstract 153 (2011).           WO         WO-2004/074290 A1 9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor Ibrutinib (PCI-32765) ha significant activity in patients with relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/005429 A1 1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of Weekly or Twice Weekly Bortezomib in Combination with								
WO         WO-2001/019829         A2         3/2001         Advani et al. Effect of Btk inhibitor PCI-32765 monotherapy on WO-2001/041754         Advani et al. Effect of Btk inhibitor PCI-32765 monotherapy on PCI-32765 monotherapy on WO-2001/044258         Advani et al. Effect of Btk inhibitor PCI-32765 monotherapy on responses in patients with relapsed aggressive NHL: Evidence of Advani et al. Preliminary Results of a Phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/038797         A2         5/2002         Annual Meeting Proceedings (Post-Meeting Edition), 28(15 MO)         WO-2002/076986         Al. 10/2002         Supp):8012 (2010).           WO         WO-2002/080926 MO         10/2002 MO         Advani et al., "The Btk inhibitor PCI-32765 is highly active and Wolden Wo-2003/00187         Wolden Wolden Wolden Molden Molde					OTHER	PUE	BLICATIONS	}
WO         WO-2001/041754 WO         6/2001 Presponses in patients with relapsed aggressive NHL: Evidence of antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/038797 A2         5/2002 Signer Ascord A		WO-2001/019829 A2						
WO         WO-2001/044258         A1         6/2001         responses in patients with relapsed aggressive NHL: Evidence of antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/038797         A2         5/2002         Annual Meeting Proceedings (Post-Meeting Edition), 28(15 Mov 2002/076986 A1 10/2002           WO         WO-2002/080926         10/2002         Supp):8012 (2010).           WO         WO-2003/000187         1/2003         Advani et al., "The Btk inhibitor PCI-32765 is highly active and well tolerated in patients (PTS) with relapsed/refractory B cell malignancies: final results from a phase I study," Ann Oncol, well tolerated in patients from a phase I study," Ann Oncol, 22(Supp 4):abstract 153 (2011).           WO         WO-2003/097645         11/2003         22(Supp 4):abstract 153 (2011).           WO         WO-2004/074290         A1         9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor Ibrutinb (PCI-32765) ha significant activity in patients with relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/005429 A1         1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of Weekly or Twice Weekly Bortezomib in Combination with				Advar	ni et al. Effect of Btl	c inhi	bitor PCI-3276	5 monotherapy on
WO         WO-2002/038797         A2         5/2002         antitumor activity from a phase I study. J. Clin. Oncol., 2010 ASCO           WO         WO-2002/076986         A1         10/2002         Supp):8012 (2010).           WO         WO-2002/080926         10/2002         Supp):8012 (2010).           WO         WO-2003/000187         1/2003         Advani et al., "The Btk inhibitor PCI-32765 is highly active and well tolerated in patients (PTS) with relapsed/refractory B cell malignancies: final results from a phase I study," Ann Oncol, wowledge well of the properties of the properti				respor	ises in patients with	relaps	sed aggressive	NHL: Evidence of
WO         WO-2002/38797         A2         5/2002         Annual Meeting Proceedings (Post-Meeting Edition), 28(15)           WO         WO-2002/076986         A1         10/2002         Supp):8012 (2010).           WO         WO-2002/080926         10/2002         Advani et al., "The Btk inhibitor PCI-32765 is highly active and well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients (PTS) with relapsed/refractory B cell malignancies: final results from a phase I study," Ann Oncol, 22(Supp 4):abstract 153 (2011).           WO         WO-2003/097645         11/2003         22(Supp 4):abstract 153 (2011).           WO         WO-2004/074290         A1         9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor Ibrutinib (PCI-32765) ha significant activity in patients with relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/005429 A1         1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of Weekly or Twice Weekly Bortezomib in Combination with				antitui	mor activity from a ph	ase I	study. J. Clin. C	Oncol., 2010 ASCO
WO         WO-2002/076986         A1         10/2002         Supp):8012 (2010).           WO         WO-2002/080926         10/2002         Advani et al., "The Btk inhibitor PCI-32765 is highly active and well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients (PTS) with relapsed/refractory B cell malignancies: final results from a phase I study," Ann Oncol, wowld-2003/097645           WO         WO-2003/097645         11/2003         22(Supp 4):abstract 153 (2011).           WO         WO-2004/074290         A1         9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor Ibrutinib (PCI-32765) ha significant activity in patients with WO-2004/100868         A2         11/2004         Ibrutinib (PCI-32765) ha significant activity in patients with relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/005429         A1         1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of Weekly or Twice Weekly Bortezomib in Combination with					•		•	
WO         WO-2002/080926         10/2002         Advani et al., "The Btk inhibitor PCI-32765 is highly active and well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients (PTS) with relapsed/refractory B cell well tolerated in patients (PTS) with relapsed/refractory B cell malignancies: final results from a phase I study," Ann Oncol, 22(Supp 4):abstract 153 (2011).           WO         WO-2003/097645         11/2003         22(Supp 4):abstract 153 (2011).           WO         WO-2004/074290 A1         9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor Ibrutinib (PCI-32765) ha significant activity in patients with relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/005429 A1         1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of Weekly or Twice Weekly Bortezomib in Combination with	WO		10/2002		~	٥	` .	,, ,,
WO         WO-2003/000187         1/2003         well tolerated in patients (PTS) with relapsed/refractory B cell           WO         WO-2003/013540         2/2003         malignancies: final results from a phase I study," Ann Oncol,           WO         WO-2003/097645         11/2003         22(Supp 4):abstract 153 (2011).           WO         WO-2004/074290         A1         9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor           WO         WO-2004/100868         A2         11/2004         Ibrutinib (PCI-32765) ha significant activity in patients with           WO         WO-2004/100868         A3         11/2004         relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/005429         A1         1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of           WO         WO-2005/014599         A1         2/2005         Weekly or Twice Weekly Bortezomib in Combination with					, ,	hibito	r PCI-32765 is	highly active and
WO         WO-2003/046200         6/2003         malignancies: final results from a phase I study," Ann Oncol,           WO         WO-2003/097645         11/2003         22(Supp 4):abstract 153 (2011).           WO         WO-2004/074290         A1         9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor           WO         WO-2004/096253         A1         11/2004         Ibrutinib (PCI-32765) ha significant activity in patients with           WO         WO-2004/100868         A2         11/2004         relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/005429         A1         1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of           WO         WO-2005/014599         A1         2/2005         Weekly or Twice Weekly Bortezomib in Combination with								~ .
WO         WO-2003/097645         11/2003         22(Supp 4):abstract 153 (2011).           WO         WO-2004/074290         A1         9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor           WO         WO-2004/096253         A1         11/2004         Ibrutinib (PCI-32765) ha significant activity in patients with           WO         WO-2004/100868         A2         11/2004         relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/005429         A1         1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of           WO         WO-2005/014599         A1         2/2005         Weekly or Twice Weekly Bortezomib in Combination with								•
WO         WO-2004/074290         A1         9/2004         Advani, R.H., et al., 2013, "Bruton tyrosine kinase inhibitor           WO         WO-2004/096253         A1         11/2004         Ibrutinib (PCI-32765) ha significant activity in patients with           WO         WO-2004/100868         A2         11/2004         relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/005429         A1         1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of           WO         WO-2005/014599         A1         2/2005         Weekly or Twice Weekly Bortezomib in Combination with				_				,
WO         WO-2004/096253         A1         11/2004         Ibrutinib (PCI-32765)         ha significant activity in patients with relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/000197         A2         1/2005         Agathocleous et al. Preliminary Results of a Phase I/II Study of Wo-2005/014599         Al 2/2005         Weekly or Twice Weekly Bortezomib in Combination with	WO						Bruton tyrosin	e kinase inhibitor
WO         WO-2004/100868 A2 WO-2004/100868 A3 I1/2004         relapsed/refractory B-cell malignancies", Journal of Clinical Oncology, vol. 31, No. 1, pp. 88-94.           WO         WO-2005/000197 A2 WO-2005/005429 A1 I/2005         1/2005 Agathocleous et al. Preliminary Results of a Phase I/II Study of Weekly or Twice Weekly Bortezomib in Combination with							•	
WO WO-2004/100868 A3 11/2004 ogy, vol. 31, No. 1 ,pp. 88-94. WO WO-2005/005429 A1 1/2005 Agathocleous et al. Preliminary Results of a Phase I/II Study of WO-2005/014599 A1 2/2005 Weekly or Twice Weekly Bortezomib in Combination with				relaps	ed/refractory B-cell m	naligna	ancies", Journa	l of Clinical Oncol-
WO WO-2005/005429 A1 1/2005 Agathocleous et al. Preliminary Results of a Phase I/II Study of Wo-2005/014599 A1 2/2005 Weekly or Twice Weekly Bortezomib in Combination with				ogy, v	ol. 31, No. 1 ,pp. 88	<b>-</b> 94.		
WO WO-2005/014599 A1 2/2005 Weekly or Twice Weekly Bortezomib in Combination with								
WO WO-2005/037836 A2 4/2005 Rituringh in Potients with Follicular Lymphoma Montle Call	WO	WO-2005/014599 A1	2/2005		•	•		
11.0 2005/05/05/07/25 72 7/2007 Kituziniao, in Taucius with Politetiai Lymphonia, Mainte Cell	WO	WO-2005/037836 A2	4/2005	Rituxi	mab, in Patients wi	th Fo	llicular Lymph	oma, Mantle Cell

#### OTHER PUBLICATIONS

Lymphoma and Waldenstrom's Macroglobulinaemia. Blood (ASH Annual Meeting Abstracts) 110:Abstract 2559 (2007).

Agency for Toxic Substances and Disease Registry, Public Health Assessment Guidance Manual, (2005).

Ahn et al. Michael acceptors as a tool for anticancer drug design. Current Pharmaceutical Design 2(3):247-262 (1996).

Apsel et al. Targeted Polypharmacology: Discovery of Dual Inhibitors of Tyrosine and Phosphoinositide Kinases. Nature Chem. Bio., 4(11):691-699 (2008).

Arnold et al. Pyrrolo[2,3-d]pyrimidines Containing an Extended 5-Substituent as Potent and Selective Inhibitors of Ick 1. Bioorg. Med. Chem. Ltrs. 10:2167-2170 (2000).

Banker et al. Modern Pharmaceutics, 3ed., Marcel Dekker, New York 1996, p. 596.

Bernstein, "Polymorphism—A Perspective," Crystal Growth & Design, 11: 632-650 (2011).

Bharate S.S., et al., "Incompatibilities of Pharmaceutical Excipients with Active Pharmaceutical Ingredients: A Comprehensive Review", Journal of Excipients and Food Chemicals, 2010, vol. 1 (3), pp. 3-26.

Biospace, 20091208, pharmacyclics, Inc. (PCYC) announces presentation of interim results from phase I trial of its first-in-human btk inhibitor PCI-32765.

Brown et al., "Phase Ib trial of AVL-292, a covalent inhibitor of Bruton's tyrosine kinase (Btk), in chronic lymphocytic leukemia (CLL) and B-non-Hodgkin lymphoma (B-NHL)," J Clin Oncol, 30(supp):abstract 8032 (2012); [online][retrieved on Jan. 14, 2016] Retrieved from the Internet: <a href="http://meetinglibrary.asco.org/content/98841-114">http://meetinglibrary.asco.org/content/98841-114</a>.

Browning. B cells move to centre stage: novel opportunities for autoimmune disease treatment. Nature Reviews/Drug Discovery 5:564-576 (Jul. 2006).

Burchat et al. Pyrazolo[3,4-d]pyrimidines Containing an Extended 3-Substituent as Potent Inhibitors of Lck—a Selectivity Insight. Bioorg. Med. Chem. Ltrs. 12:1687-1690 (2002).

Burger et al. CXCR4 antagonists: targeting the microenvironment in leukemia and other cancers, Leukemia 23:43-52 (2009).

Burger et al. High-Level Expression of the T-Cell Chemokines CCL3 and CCL4 by Chronic Lymphocytic Leukemia B Cells in Nurselike Cell Cocultures and After BCR Stimulation. Blood 113(13):3050-3058 (2008).

Burger. Targeting the microenvironment in chronic lymphocytic leukemia is changing the therapeutic landscape. Curr. Opin. Oncol. 24(6):643-649 (Epub Sep. 6, 2012/Nov. 2012).

Byrd et al. Entering the era of targeted therapy for chronic lymphocytic leukemia: impact on the practicing clinician. J. Clinical Oncology (Jul. 21, 2014) (pii: JCO.2014.55.8262).

Byrd et al. Targeting BTK with Ibrutinib in relapsed chronic lymphocytic leukemia, NEJM 369(1):32-42 (Jul. 4, 2013).

Byrd J.C., et al., "Three-year follow-up of treatment-naïve and previously treated patients with CLL and SLL receiving single-agent ibrutinib", Blood, Apr. 16, 2015, vol. 125 (16), pp. 2497-2506. Byrn et al., "Pharmaceutical Solids: A Strategic Approach to Regulatory Considerations," Pharmaceutical Research, 12: 945-954 (1995).

Calderwood et al., "Pyrrolo[2,3-d]pyrimidines Containing Diverse N-7 Substituents as Potent Inhibitors of Lcks," Bioorg Med Chem Lett, 12: 1683-1686 (2002).

Cameron F., et al., "Ibrutinib: first global approval," Drugs, Feb. 2014, vol. 74 (2), pp. 263-271.

Cannon. Chapter Nineteen in Burger's Medicinal Chemistry and Drug Discovery, Fifth Edition, vol. 1: Principles and Practice, Wiley-Interscience 1995, pp. 783-802.

Carmi et al. Clinical perspectives for irreversible tyrosine kinase inhibitors in cancer. Biochem. Pharmacol. (Epub Aug. 4, 2012) 84(11):1388-1399 (Dec. 2012).

Carrle et al. Current Strategies of Chemotherapy in Osteosarcoma. International Orthopaedics 30:445-451 (2006).

Certified Translation of CN103121999A, dated Sep. 29, 2016.

Certified Translation of CN103923084A, dated Sep. 29, 2016.

Certified Translation of CN103923084B, dated Sep. 29, 2016.

Certified Translation of CN104523695A, dated Nov. 14, 2016.

Chang et al. Egress of CD19+CD5+ cells into peripheral blood following treatment with the Bruton tyrosine kinase inhibitor inbrutinib in mantel cell lymphoma patients. Blood, 122:2412-2424 (2013).

Chang et al., "PCI-45292, a novel Btk inhibitor with optimized pharmaceutical properties, demonstrates potent activities in rodent models of arthritis," ACR/ARHP Scientific Meeting, Poster #286 (2010).

Chang et al., "The Bruton tyrosine kinase inhibitor PCI-32765 ameliorates autoimmune arthritis by inhibition of multiple effector cells," Arhtritis Res Ther, 13:R115 (2011).

Chavez et al. Ibrutinib: An Evidence-Based Review of Its Potential in the Treatment of Advanced Chronic Lymphocytic Leukemia. Core Evidence 8:37-45 (2013).

Chen et al. SYK-dependent tonic B-cell receptor signaling is a rational treatment target in diffuse large B-cell lymphoma. Blood 111(4):2230-2237 (2008) [E-pub Nov. 15, 2007].

Ciric et al., "Clonal Evolution in Waldenstrom macroglobulinemia Highlights Functional Role of B-Cell Receptor," Blood, 97: 321-323 (2001)

Cohen et al. Structural Bioinformatics-Based Design of Selective, Irreversible Kinase Inhibitors. Science 308:1318-1321 (May 27, 2005).

Czuczman et al. Rituximab in combination with fludarabine chemotherapy in low-grade or follicular lymphoma. J. Clin. Oncol. 23(4):694-704 (Feb. 1, 2005).

D'Cruz et al. Novel Bruton's tyrosine kinase inhibitors currently in development. OncoTargets and Therapy 6:161-176 (2013).

Dana-Farber Cancer Institute. A Phase II Study of Ibrutinib Plus FCR in Previously Untreated, Younger Patients With Chronic Lymphocytic Leukemia (iFCR). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Sep. 23, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02251548?term=NCT02251548 NLM Identifier: NCT02251548.

Dana-Farber Cancer Institute. Ibrutinib (PCI-32765) in Waldenstrom's Macroglobulinemia. In: Clinical Trials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). May 17, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01614821 NLM Identifier: NCT01614821.

Davids et al. Targeting the B Cell Receptor Pathway in Chronic Lymphocytic Leukemia. Leuk. Lymphoma (Epub May 23, 2012), 53(12):2362-2370 (Dec. 2012).

Davis et al., "Chronic active B-cell receptor signalling in diffuse large B-cell lymphoma," Nature, 463(7277):88-92 (2010).

Desiderio. Role of Btk in B cell development and signaling. Curr. Op. in Immunology 1997, 9:534-540.

Devos et al., "The Bruton's tyrosine kinase (BTK) inhibitor, ibrutinib (PCI-32765), has preferential activity in the activated B cell-like (ABC) subtype of relapsed/refractory (RR) DLBCL: interim phase 2 results," Haematologica 98(s1):490 (2013).

Dias et al. Ibrutinib: A New Frontier in the Treatment of Chronic Lymphocytic Leukemia by Bruton's Tyrosine Kinase Inhibition. Cardiovascular & Hematological Agents in Medicinal Chemistry 11:265-271 (2013).

Dorwald, F.Z., "Side Reactions in Organic Synthesis," Wiley:VCH, Weinheim p. IX of Preface, Wiley-VCH Verlag GmbH & C KGaA (2005).

EA200901313 Notification of Office Action dated Oct. 31, 2011. EA201000599 Search Report dated Nov. 15, 2010.

Edwards. BTK inhibition in myeloma: targeting the seed and the soil. Blood 120(9):1757-1759 (Aug. 2012).

EP 06850039.6 Search Report and Written Opinion dated Feb. 15, 2010

 $\rm EP~06850386.1~Search~Report~and~Written~Opinion~dated~Sep.~10,~2010.$ 

EP 08744513.6 Examination Report dated Jan. 16, 2013.

EP 08744513.6 Search Report and Written Opinion dated Mar. 18, 2010.

#### OTHER PUBLICATIONS

EP 09798770.5 Search Report and Written Opinion dated Oct. 28, 2011

EP 10155834.4 Search Report and Written Opinion dated May 27, 2010

EP 10823966 Supplementary European Search Report dated Oct. 17, 2011.

EP 10823966.6 Search Report dated Oct. 17, 2011.

EP 10823966.6 Written Opinion dated Dec. 6, 2011.

EP 12151943.3 Examination Report dated Feb. 5, 2013.

EP 12151943.3 Search Report and Written Opinion dated Mar. 13, 2012

EP 12166295.1 Search Report and Written Opinion dated Nov. 6, 2012.

EP 12166296.9 Search Report and Written Opinion dated Nov. 8, 2012.

EP 12166298.5 Search Report and Written Opinion dated Nov. 7, 2012.

EP 12166300.9 Search Report and Written Opinion dated Oct. 31, 2012

EP 12166301.7 Search Report and Written Opinion dated Nov. 6, 2012

EP 12166302.5 Search Report and Written Opinion dated Nov. 6, 2012.

EP 12166305.8 Examination Report dated Dec. 3, 2013.

EP 12166305.8 Search Report and Written Opinion dated Nov. 6, 2012

EP 12166306.6 Search Report and Written Opinion dated Nov. 8, 2012

EP 12172840.6 Search Report and Written Opinion dated Dec. 12, 2012.

EP 12172841.4 Search Report and Written Opinion dated Jan. 2, 2013.

EP 12172842.2 Extended Search Report dated May 14, 2013.

EP 12172842.2 Partial Search Report dated Jan. 24, 2013.

EP 12172843.0 Search Report and Written Opinion dated Jan. 18, 2013.

EP06850039 Supplemental Search Report dated Feb. 15, 2010.

EP06850039 Supplemental Search Report dated Feb. 9, 2010.

EP11790524.0 Extended European Search Report dated Oct. 9, 2013

EP12172840.6 Office Action dated Jun. 12, 2014.

EP12172841.4 Office action dated Jun. 12, 2014.

EP12172842.2 Office Action dated Jul. 1, 2014.

EP12172843.0 Office action dated Jul. 1, 2014.

EP15170739.5 Extended European Search Report, dated Nov. 10, 2015

Fabian et al. A small molecule-kinase interaction map for clinical kinase inhibitors. Nature Biotechnology, 23(3): 329-336 (2005).

FDA, International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH), Federal Register of Nov. 25, 1997, draft of "Q6A Specifications: Test Procedures and Acceptance Criteria for New Drug Substances and New Drug Products: Chemical Substances." endorsed final guidance Oct. 6, 1999.

Federal Drug Adminstration, Draft Guidance for Industry, ANDAs: Pharmaceutical Solid Polymorphism at 3 (Dec. 2004).

Fedorak et al. A novel colon-specific steroid prodrug enhances sodium chloride absorption in rat colitis. Am. J. Physiol. 269:G210-218 (1995).

Fisher et al. Prolonged disease-free survival in Hodgkin's disease with MOPP reinduction after first relapse. Ann. Intern. Med., 90(5):761-763 (1979).

Fowler et al., "The Bruton's tyrosine kinase inhibitor ibrutinib (PCI-32765) is active and tolerated in relapsed follicular lymphoma," 54th American Society of Hematology Annual Meeting and Exposition, Abstract 156 (2012).

Friedberg et al. Inhibition of Syk with fostamatinib disodium has significant clinical activity in non-Hodgkin lymphoma and chronic lymphocytic leukemia. Blood 115(13):2578-2585 (2010) [E-pub Nov. 17, 2009].

Fruman. Xid-like Phenotypes: A B Cell Signalosome Takes Shape. Immunity 13:1-3 (Jul. 2000).

Fry et al., "Specific Irreversible Inactivation of the Epidermal Growth Factor Receptor and erbB2, by a New Class of Tyrosine Kinase Inhibitor," Proc Natl Acad Sci, 95: 12022-12027 (1998). Gazitt et al. Differential mobilization of CD34+ Cells and lymphoma cells in non-Hodgkin's lymphoma patients mobilized with

phoma cells in non-Hodgkin's lymphoma patients mobilized with different growth factors, J of Hematotherapy & Stem Cell Research 10:167-176 (2001).

Ghia. Ibrutinib: better combined with other drugs? Lancet 15:1043-1044 (2014).

Giuliani. Multiple myeloma bone disease: pathophysiology of osteoblast inhibition. Blood (Epub Aug. 17, 2006) 108(13):3992-3996 (2006).

Glassman et al., "The value of fluorescence in situ hybridization in the diagnosis and prognosis of chronic lymphocytic leukemia," Cancer Genet Cytogen, 158:88-91 (2005).

Gold. To make antibodies or not: signaling by the B-cell antigen receptor. Trends in Pharmacological Sciences, 23(7):316-324 (Jul. 2002)

Gordon et al. Somatic hypermutation of the B cell receptor genes B29 (Igb, CD79b) and mb1 (Iga, CD79a). PNAS 100(7):4126-4131 (2003)

Groshock et al. Molecular Target Class is Predictive of In vitro Response Profile. Cancer Res. 70:3677-3686 (2010).

Gu et al., "Polymorph Screening: Influence of Solvent on the Rate of Solvent-Mediated Polymorphic Transformation," Journal of Pharmaceutical Sciences 90, 11: 1878-1890 (2001).

Hagemeister. Rituximab for the treatment of non-Hodgkin's lymphoma and chronic lymphocytic leukaemia. Drugs 70(3):261-272 (2010).

Haleblian and McCrone, "Pharmaceutical Applications of Polymorphism," 58(8): 911-929 (1969).

Hantschel et al. The Btk Tyrosine Kinase is a Major Target of the Bcr-Ab1 Inhibitor Dasatinib. PNAS 104(33):13283-13288 (2007). Hata et al. Bruton's tyrosine kinase-mediated Interleukin-2 gene activation in mast cells. J. Biol. Chem. 273(18): 10979-10987 (1998).

Herman et al. Bruton tyrosine kinase represents a promising therapeutic target for treatment of chronic lymphocytic leukemia and is effectively targeted by PCI-32765. Blood (Epub Mar. 21, 2011), 117(23):6287-6296 (Jun. 2011).

Hiddeman et al. Rituximab Plus Chemotherapy in Follicular and Mantle Cell Lymphomas, Seminars in Oncology 30(1)Suppl.2:16-20 (Feb. 2003).

Hiddeman et al., "Frontline therapy with rituximab added to the combination of cyclophosphamide, doxorubicin, vincristine, and prednisone (CHOP) significantly improves the outcome for patients with advanced-stage follicular lymphoma compared with therapy with CHOP alone: results of a prospective randomized study of the German low-grade lymphoma study group," Blood, 106(12):3725-32 (2005).

Higuchi et al. Pro-drugs as Novel Delivery Systems, vol. 14 of the A.C.S. Symposium Series (1975).

Hochhaus et al. A selective HPLC/RIA for dexamethasone and its prodrug dexamethasone-21-sulphobenzoate sodium in biological fluids. Biomed. Chrom., 6:283-286 (1992).

Honigberg et al. Targeting Btk in lymphoma: PCI-32765 inhibits tumor growth in mouse lymphoma models and a fluorescent analog of PCI-32765 is an active-site probe that enables assessment of Btk inhibition in vivo. ASH Annual Meeting Abstracts 1592. 110(11): 475A (2007).

Honigberg et al., "The Bruton Tyrosine Kinase Inhibitor PCI-32765 Blocks B-Cell Activation and is Efficacious in Models of Autoimmune Disease and B-Cell Malignancy," PNAS USA, 107: 13075-13080 (2010).

Horwood et al. Bruton's Tyrosine Kinase is Required for Lipopolysaccharide-induced Tumor Necrosis Factor a Production. J. Exp. Med. 197(12):1603-1611 (Jun. 2003).

#### OTHER PUBLICATIONS

http://www.uspto.gov/web/offices/pac/dapp/1pecba.htm#7, last accessed Feb. 16, 2011.

Huhn et al. Rituximab therapy of patients with B-cell chronic lymphocytic leukemia. Blood 98(5):1326-1331 (Sep. 1, 2001).

International Search Report and Written Opinion for Application No. PCT/US2016/020467, dated May 23, 2016, 10 pages.

International Search Report for International Application No. PCT/US2006/049626 dated Apr. 9, 2008.

International Search Report for International Application No. PCT/US2009/050897 dated Mar. 15, 2010.

International Search Report for International Application No. PCT/US2016/024305 dated Jun. 27, 2016.

International Search Report for International Application No. PCT/US2016/024321 dated Jun. 28, 2016.

Iqbal et al., on pp. 2-4 (Molecular Biology International, 2014, Article ID 852748, 9 pages.

Iwaki et al. Btk Plays a Crucial Role in the Amplification of FceRI-mediated Mast Cell Activation by Kit. J. Biol. Chem. 280(48):40261-40270 (Dec. 2, 2005).

Jaffe. The 2008 WHO classification of lymphomas: implications for clinical practice and translational research. Hematology 1:523-531 (2009).

Janssen Biotech, Inc. An open label treatment use protocol for ibrutinib in subjects with relapsed or refractory mantel cell lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Apr. 6, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01833039 NLM Identifier: NCT01833039.

Janssen Pharmaceutical K.K. A study to evaluate the Bruton's tyrosine kinase (Btk) inhibitor PCI-32765 in patients with recurrent mature B-cell neoplasms. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 9, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01704963 NLM Identifier: NCT01704963.

Janssen Pharmaceutical K.K. Study of the Bruton's Tyrosine Kinase (BTK) Inhibitor Ibrutinib in Participants With Relapsed or Refractory Mantle Cell Lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jun. 19, 2014—[cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02169180?term=NCT02169180 NLM Identifier: NCT02169180.

Janssen Research & Development, LLC. A Study to Evaluate the Effects of Ibrutinib on Cardiac Repolarization in Healthy Participants. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 20, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02271438?term=NCT02271438 NLM Identifier: NCT02271438.

Janssen Research & Development, LLC. Pharmacokinetic and Pharmacodynamic Study to Evaluate Safety and Efficacy of the Combination of Ibrutinib With Nivolumab in Participants With Hematologic Malignancies. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Dec. 30, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02329847?term=NCT02329847 NLM Identifier: NCT02329847.

Janssen Research and Development, LLC. A long-term extension study of PCI-32765 (Ibrutinib). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Mar. 4, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01804686 NLM Identifier: NCT01804686.

Janssen Research and Development, LLC. A pharmacokinetic study in healthy participants to assess the pharmacokinetics and safety of a supratherapeutic dose of PCI-32765 (Ibrutinib) capsule and solution formulations administered with food. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Aug. 19, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01969266 NLM Identifier: NCT01969266.

Janssen Research and Development, LLC. A study combining Ibrutinib with rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone in patients with CD20-positive B-cell non Hodgkin lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Mar. 30, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01569750 NLM Identifier: NCT01569750.

Janssen Research and Development, LLC. A study of ibrutinib in combination with bendamustine and rituximab in patients with relapsed or refractory chronic lymphocytic leukemia or small lymphocytic lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). May 15, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01611090 NLM Identifier: NCT01611090.

Janssen Research and Development, LLC. A study of PCI-32765 (Ibrutinib) in combination with either bendamustine and rituximab or rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone in patients with previously treated indolent non-Hodgkin lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 28, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01974440 NLM Identifier: NCT01974440.

Janssen Research and Development, LLC. A study of Pci-32765 (Ibrutinib) in patients with refractory follicular lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jan. 25, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01779791 NLM Identifier: NCT01779791.

Janssen Research and Development, LLC. A study of the Bruton's tyrosine kinase inhibitor ibrutinib given in combination with bendamustine and rituximab in patients with newly diagnosed mantel cell lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jan. 24, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01776840 NLM Identifier: NCT01776840.

Janssen Research and Development, LLC. A study of the Bruton's tyrosine kinase inhibitor PCI-32765 (Ibrutinib) versus rituximab in patients with relapsed or refractory chronic lymphocytic leukemia/small lymphocytic lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 25, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01973387 NLM Identifier: NCT01973387.

Janssen Research and Development, LLC. A study on the Bruton's tyrosine kinase inhibitor, PCI-32765 (Ibrutinib), in combination with rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone in patients with newly diagnosed non-germinal center B-cell subtype of diffuse large B-cell lymphoma. In: ClinicalTrials. gov [Internet]. Bethesda (MD): National Library of Medicine (US). May 14, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01855750 NLM Identifier: NCT01855750.

Janssen Research and Development, LLC. A study to assess the absolute bioavailability of Oral PCI-32765 and the effect of grape-fruit juice on the bioavailability of PCI-32765 in healthy participants. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). May 28, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/study/NCT01866033 NLM Identifier: NCT01866033.

Janssen Research and Development, LLC. A study to assess the effect of ketoconazole on the pharmacokinetics of ibrutinib in healthy participants. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jun. 18, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01626651 NLM Identifier: NCT01626651.

Janssen Research and Development, LLC. A study to assess the effect of rifampin on the pharmacokinetics of PCI-32765 in healthy participants. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jan. 4, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01763021 NLM Identifier: NCT01763021.

Janssen Research and Development, LLC. A study to determine the absorption, metabolism, and routes of excretion of (14C) radiolabeled ibrutinib in healthy male participants. In: ClinicalTri-

#### OTHER PUBLICATIONS

als.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Aug. 9, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01674322 NLM Identifier: NCT01674322.

Janssen Research and Development, LLC. A study to determine the effect of food on the pharmacokinetics of PCI-32765. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Mar. 4, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01820936 NLM Identifier: NCT01820936.

Janssen Research and Development, LLC. A study to evaluate the efficacy and safety of ibrutinib, in patients with mantel cell lymphoma who progress after bortezomib therapy. In: ClinicalTrials. gov [Internet]. Bethesda (MD): National Library of Medicine (US). May 14, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01599949 NLM Identifier: NCT01599949.

Janssen Research and Development, LLC. A study to evaluate the pharmacokinetics of PCI-32765 in participants with varying degrees of hepatic impairment. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jan. 9, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01767948 NLM Identifier: NCT01767948.

Janssen Research and Development, LLC. Study of ibrutinib (a Bruton's tyrosine kinase inhibitor), versus temsirolimus in patients with relapsed or refractory mantel cell lymphoma who have received at least one prior therapy. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jul. 18, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01646021 NLM Identifier: NCT01646021.

Jefferies et al., "Bruton's tyrosine kinase is a toll/interleukin-1 receptor domain-binding protein that participates in nuclear factor kB activation by toll-like receptor 4," J Biol Chem, 278:26258-64 (2003).

Kamb. What's wrong with our cancer models? Nature Reviews Drug Discovery 4:161-165 (2005).

Kawakami et al. Terreic acid, a quinone epoxide inhibitor of Bruton's tyrosine kinase. PNAS USA 96:2227-2232 (1999). Korade-Mirnics et al. Src kinase-mediated signaling in leukocytes.

Korade-Mirnics et al. Src kinase-mediated signaling in leukocytes J. Leukoc. Bio., 68(5):603-613 (Nov. 2000).

Kozaki et al. Development of a Bruton's tyrosine kinase (Btk) inhibitor -ONO-WG-307, a potential treatment for B-cell malignancies. 53rd American Society of Hematology Annual Meeting and Exposition, San Diego, CA, Poster #857 (Dec. 10-13, 2011). Kuglstatter et al. Insights into the conformational flexibility of Bruton's tyrosine kinase from multiple ligand complex structures. Protein Science 20(2):428-436 (2011) [E-pub Dec. 17, 2010].

Kuppers. Mechanisms of B-cell lymphoma pathogenesis. Nature Reviews/Cancer 5:251-262 (2005).

Kurosaki. Functional dissection of BCR signaling pathways. Curr. Op. Imm. 12:276-281 (2000).

Kushner et al. Pharmacological uses and perspective of heavy water and deuterated compounds. Canadian Journal of Physiology and Pharmacology 77(2):79-88 (1999).

Label for TASIGNA® (nilotinib), revised Aug. 2009.

Larsen et al., "Prodrug Forms for the Sulfonamide Group. II. Water-soluble Amino Acid Derivatives of N-methylsulfonamides as Possible Prodrugs," Int J Pharmaceutics, 47: 103-110 (1988).

Larsen et al. Prodrug forms for the sulfonamide group. I. Evaluation of N-acyl derivatives, N-sulfonylamindes, N-sulfonylsulfilimines and sulfonylureas as possible prodrug derivatives. Int. J. Pharmaceutics 37:87-95 (1987).

Le Tourneau et al. Dose Escalation Methods in Phase I Cancer Clinical Trials. J. Natl Cancer 101:708-720 (2009).

Li et al. Activation of Bruton's Tyrosine Kinase (BTK) by a Point Mutation in its Pleckstrin Homology (PH) domain. Immunity 2:451-460 (1995).

Lim et al. Asymmetric syntheses of fused bicyclic lactams. Journal of Organic chemistry 66(26):9056-9062 (2001).

Lin et al. Selective Itk inhibitors block T-cell activation and murine lung inflammation, Biochemistry 43:11056-11062 (2004).

Liu et al. Structural Basis for selective inhibition of Src family kinases by PPI. Chemistry and Biology 6:671-678, in particular table 1, p. 671 (1999).

Lossos. Molecular Pathogenesis of Diffuse Large B-Cell Lymphoma. J. Clin. Oncol. 23(26):6351-6357 (Sep. 10, 2005).

Lou et al. Bruton's tyrosine kinase inhibitors: approaches to potent and selective inhibition, preclinical and clinical evaluation for inflammatory diseases and B cell malignancies. J Med Chem. May 24, 2012;55(10):4539-50 Publication Date (Web): Mar. 6, 2012. Luskova et al. Modulation of the Fce Receptor I Signaling by Tyrosine Kinase Inhibitors: Search for Therapeutic Targets of Inflammatory and Allergy Diseases. Curr. Pharmaceutical Design 10:1727-1737 (2004).

M.D. Anderson Cancer Center. A Phase I/II Study of Ibrutinib in Previously Treated Epidermal Growth Factor Receptor (EGFR) Mutant Non-Small Cell Lung Cancer. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Dec. 17, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02321540?term=NCT02321540 NLM Identifier: NCT02321540.

M.D. Anderson Cancer Center. A Phase I/II Trial of PCI-32765 (BTK Inhibitor) in Combination With Carfilzomib in Relapse/Refractory Mantle Cell Lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 16, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02269085?term=NCT02269085 NLM Identifier: NCT02269085.

M.D. Anderson Cancer Center. Ibrutinib Post Stem Cell Transplantation (SCT) in Double-Hit B-Cell Lymphoma. In: ClinicalTrials. gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 21, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02272686?term=NCT02272686 NLM Identifier: NCT02272686.

M.D. Anderson Cancer Center. Ibrutinib versus ibrutinib + rituximab (i vs iR) in patients with relapsed chronic lymphocytic leukemia (CLL). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Dec. 5, 2013—[cited Apr. 15, 2014]. Available from: http://clinicaltrials.gov/ct2/show/NCT02007044 NLM Identifier: NCT02007044.

M.D. Anderson Cancer Center. Phase 2 ibrutinib + rituximab in relapsed/refractory mantel cell lymphoma (R/R MCL). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jun. 14, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01880567 NLM Identifier: NCT01880567.

M.D. Anderson Cancer Center. Phase 2 study of the combination of Bruton's tyrosine kinase inhibitor PCI-32765 and rituximab in high-risk chronic lymphocytic leukemia and small lymphocytic lymphoma patients. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jan. 25, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01520519 NLM Identifier: NCT01520519.

M.D. Anderson Cancer Center. Pilot study to determine effects of the Btk inhibitor PCI-32765 on leukemia cell kinetics and trafficking, using heavy water labeling in subjects with CLL and SLL. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Dec. 13, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01752426 NLM Identifier: NCT01752426.

MacPartlin et al. Bruton's tyrosine kinase is not essential for Bcr-Abl-mediated transformation of lymphoid or myeloid cells. Leukemia 22:1354-1360 (2008).

Maddocks et al. Ibrutinib in B-cell lymphomas. Current Treatment Options in Oncology 15:226-237 (2014) (Epub: Feb. 1, 2014).

Mahajan et al. Rational Design and Synthesis of a Novel Antileukemic Agent Targeting Bruton's Tyrosine Kinase (BTK), LFM-A13 [a-Cyano-β-Methyl-N-(2,5-Dibromophenyl)Propenamide]. J. of Biol. Chem. 274(14):9587-9599 (1999).

Mallis et al. Structural characterization of a proline-driven conformational switch within the Itk SH2 domain. Nat. Struct. Biol., 9(12):900-905 (2002).

#### OTHER PUBLICATIONS

Mangla et al., "Pleiotropic consequences of Bruton tyrosine kinase deficiency in myeloid lineages lead to poor inflammatory responses," Blood, 104(4):1191-1197 (2004).

Marina et al. Biology and Therapeutic Advances for Pediatric Osteosarcoma. The Oncologist 9:422-441 (2004).

McConathy et al. Stereochemistry in Drug Action. J Clinical Psychiatry. 5:70-73 (2003).

McLeod et al., "A glucocorticoid prodrug facilitates normal mucosal function in rat colitis without adrenal suppression," Gastroenterol, 106:405-413 (1994).

Memorial Sloan-Kettering Cancer Center. Bruton's Tyrosine Kinase (BTK) Inhibitor, Ibrutinib, in Patients With Refractory/Recurrent Primary Central Nervous System Lymphoma (PCNSL) and Refractory/Recurrent Secondary Central Nervous System Lymphoma (SCNSL). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Dec. 9, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02315326?term=NCT02315326 NLM Identifier: NCT02315326.

Mendel et al., "In vivo Antitumor Activity of SU11248, A Novel Tyrosine Kinase Inhibitor Targeting Vascular Endothelial Growth Factor and Platelet-derived Growth Factor Receptors: Determination of a Pharmacokinetic/pharmacodynamic Relationship," Clin Cancer Res, 9(1): 327-337 (2003).

Merged Markush Service Search, Jun. 27, 2005.

Middendorp et al. Function of Bruton's Tyrosine Kinase during B Cell Development is Partially Independent of its Catalytic Activity. J Immunol 171:5988-5996 (2003).

Middendorp et al. Tumor Suppressor Function of Bruton Tyrosine Kinase is Independent of its catalytic activity. Blood 105(1):259-261 (2005).

Morissette et al., "High-throughput Crystallization: Polymorphs, Salts, Co-Crystals and Solvates of Pharmaceutical Solids," Advanced Drug Delivery Review, 56: 275-300 (2004).

Mukoyama et al., "Preparation of imidazol [1,5-a]pyrazine derivatives, pharmaceutical compositions containing them, and their uses for prevention or treatment of protein tyrosine kinase-related diseases," retrieved from STN Database Accession No. 2005:299462 Patent No. JP2005089352, Abstract (2005).

National Cancer Institute (NCI). A multicenter phase 2 study of the Bruton's tyrosine kinase inhibitor PCI-32765 for treatment of relapsed hairy cell leukemia. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Nov. 2, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01981512 NLM Identifier: NCT01981512.

National Cancer Institute (NCI). Ibrutinib and rituximab compared with fludarabine phosphate, cyclophosphamide, and rituximab in treating patients with untreated chronic lymphocytic leukemia or small lymphocytic lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jan. 27, 2014—[cited Apr. 15, 2014]. Available from: http://clinicaltrials.gov/ct2/show/NCT02048813 NLM Identifier: NCT02048813.

National Cancer Institute (NCI). Ibrutinib in treating patients with relapsed hairy cell leukemia. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Apr. 24, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01841723 NLM Identifier: NCT01841723.

National Cancer Institute (NCI). Ibrutinib in treating patients with relapsed or refractory follicular lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). May 6, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01849263 NLM Identifier: NCT01849263.

National Cancer Institute (NCI). Lenalidomide and ibrutinib in treating patients with relapsed or refractory B-Cell non-Hodgkin lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Sep. 27, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01955499 NLM Identifier: NCT01955499.

National Cancer Institute (NCI). Rituximab and bendamustine hydrochloride, rituximab and ibrutinib, or ibrutinib alone in treating older patients with previously untreated chronic lymphocytic leukemia. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jun. 24, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01886872 NLM Identifier: NCT01886872.

National Cancer Institute (NCI). Rituximab, lenalidomide, and ibrutinib in treating patients with previously untreated stage II-IV follicular lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Apr. 9, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01829568 NLM Identifier: NCT01829568.

National Cancer Institute. Ibrutinib and Combination Chemotherapy in Treating Patients With Relapsed or Refractory Diffuse Large B-Cell Lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jul. 16, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02219737?term=NCT02219737 NLM Identifier: NCT02219737.

National Cancer Institute. Ibrutinib and Palbociclib Isethionate in Treating Patients With Previously Treated Mantle Cell Lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jun. 9, 2014—[cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02159755?term=NCT02159755 NLM Identifier: NCT02159755.

National Cancer Institute. Ibrutinib in Treating Patients With Relapsed or Refractory B-cell Acute Lymphoblastic Leukemia. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Apr. 30, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/

NCT02129062?term=NCT02129062 NLM Identifier: NCT02129062.

National Cancer Institute. Ibrutinib in Treating Relapsed or Refractory B-cell Non-Hodgkin Lymphoma in Patients With HIV infection. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Apr. 7, 2014 [ cited Feb. 5, 2015]. Available from: https://clinicaltrial.gov/ct2/show/NCT02109224?term=NCT02109224. NLM Identifier: NCT02109224.

National Cancer Institute. Lenalidomide, Ibrutinib, and Rituximab in Treating Patients With Relapsed or Refractory Chronic Lymphocytic Leukemia or Small Lymphocytic Lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). May 30, 2014 [cited Feb. 15, 2015] Available from: https://clinicaltrial.gov/ct2/show/

NCT02160015?term=NCT02160015 NLM Identifier: NCT02160015.

National Cancer Institute. Phase 1 Study of Ibrutinib and Immuno-Chemotherapy Using Dose-Adjusted-Temozolomide, Etoposide, Doxil, Dexamethasone, Ibrutinib,Rituximab (DA-TEDDI-R) in Primary CNS Lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jul. 29, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02203526?term=NCT02203526 NLM Identifier: NCT02203526.

National Center Institute (NCI). Lenalidomide and Ibrutinib in treating patients with relapsed or refractory chronic lymphocytic leukemia or small lymphocytic lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jun. 24, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01886859 NLM Identifier: NCT01886859. National Heart, Lung, and Blood Institute (NHLBI). PCI-32765 for special cases of chronic lymphocytic leukemia or small lymphocytic lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Dec. 22, 2011—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01500733 NLM Identifier: NCT01500733.

Niiro et al. Regulation of B-Cell Fate by Antigen-Receptor Signals. Nature Reviews 2:945-956 (2002).

#### OTHER PUBLICATIONS

Nisitani et al. In situ detection of activated Bruton's tyrosine kinase in the Ig signaling complex by phosphopeptide-specific monoclonal antibodies. PNAS USA 96:2221-2226 (1999).

Nogrady (1985) Medicinal Chemistry a Biochemical Approach, Oxford University Press, New York, pp. 388-394.

Northwestern University. Ibrutinib After Intensive Induction in Treating Patients With Previously Untreated Mantle Cell Lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Sep. 12, 2014 [cited 2-15 Feb. 5,] Available from: https://clinicaltrial.gov/ct2/show/NCT02242097?term=NCT02242097 NLM Identifier: NCT02242097.

Ohio State University Comprehensive Cancer Center. PCI-32765 (Ibrutinib) in treating patients with relapsed or refractory chronic lymphocytic leukemia, small lymphocytic lymphoma, or B-cell prolymphocytic leukemia. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Apr. 23, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01589302 NLM Identifier: NCT01589302.

Ohio State University Comprehensive Cancer Center. Rituxan/Bendamustine/PCI-32765 in relapsed DLBCL, MCL, or indolent non-Hodgkin's lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Nov. 1, 2011—[cited Feb. 6, 2014]. Available from: http://clinicaltrials.gov/ct2/show/NCT01479842 NLM Identifier: NCT01479842.

Oligino et al. Targeting B cells for the treatment of rheumatoid arthritis. Arthritis Res. Ther., 5(Suppl.4):S7-S11 (2002).

Pagel et al., "Induction of apoptosis using inhibitors of lysophosphatidic acid acyltransferase-beta and anti-CD20 monoclonal antibodies for treatment of human non-Hodgkin's lymphomas," Clin Cancer Res, 11(13):4857-4866 (2005).

Pan et al., "Discovery of selective irreversible inhibitors for Bruton's tyrosine kinase," ChemMedChem, 2:58-61 (2007).

PCT/US06/49626 Search Report dated Apr. 9, 2008.

PCT/US08/058528 Search Report and Written Opinion dated Sep. 30, 2008.

PCT/US09/50897 IPER and Written Opinion dated Jan. 27, 2011. PCT/US09/50897 Search Report dated Mar. 15, 2010.

PCT/US10/52377 Search Report and Written Opinion dated Jun. 29, 2011

PCT/US2006/49626 International Preliminary Report on Patentability Search Report dated Mar. 24, 2009.

PCT/US2008/058528 International Preliminary Report on Patent-

ability Search Report dated Sep. 29, 2009. PCT/US2008/058528 International Search Report and Written

Opinion dated Sep. 30, 2008. PCT/US2009/50897 International Preliminary Examination Report

and Written Opinion dated Jan. 18, 2011.

PCT/US2010/52377 International Search Report and Written Opinion dated Jun. 29, 2011.

PCT/US2013/043888 International Preliminary Report on Patentability dated Dec. 9, 2014.

PCT/US2013/043888 International Search Report and Written Opinion dated Sep. 23, 2013.

PCT/US2015/044258 International Search Report and Written Opinion dated Nov. 6, 2015.

PCT/US2015/16895 International Search Report and Written Opinion dated May 22, 2015.

PCT/US2015/21871 International Search Report and Written Opinion dated Jul. 8, 2015.

Peterson et al. Prolonged single-agent versus combination chemotherapy in indolent follicular lymphomas: a study of the cancer and leukemia group. Br. J. Clin. Oncol., 21(1):5-15 (Jan. 1, 2003).

Pharmacyclics, Inc. A Multi-Center Study of Ibrutinib in Combination With Obinutuzumab Versus Chlorambucil in Combination With Obinutuzumab in Patients With Treatment naïve CLL or SLL. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 1, 2014 [cited Feb. 5, 2015] Available from:

https://clinicaltrial.gov/ct2/show/

NCT02264574?term=NCT02264574 NLM Identifier: NCT02264574.

Pharmacyclics, Inc. A multicenter phase 2 study of PCI-32765 (Ibrutinib) in patients with relapsed or refractory chronic lymphocytic leukemia (CLL) or small lymphocytic lymphoma (SLL) with 17p deletion. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Dec. 3, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01744691 NLM Identifier: NCT01744691.

Pharmacyclics, Inc. A multicenter, open-label, phase 3 study of the Bruton's tyrosine kinase inhibitor PCI-32765 versus chlorambucil in patients 65 years or older with treatment-naive chronic lymphocytic leukemia or small lymphocytic lymphoma (RESONATE-2). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 29, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01722487 NLM Identifier: NCT01722487.

Pharmacyclics, Inc. A phase 3 study of ibrutinib (PCI-32765) versus of atumumab in patients with relapsed or refractory chronic lymphocytic leukemia (RESONATE). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Apr. 11, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01578707 NLM Identifier: NCT01578707.

Pharmacyclics, Inc. An open-label extension study in patients 65 years or older with chronic lymphocytic leukemia (CLL) or small lymphocytic lymphoma (SLL) who participated in study PCYC-115-CA (PCI-32765 versus chlorambucil). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Nov. 2, 2012—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01724346 NLM Identifier: NCT01724346.

Pharmacyclics, Inc. Efficacy and safety study of PCI-32765 combined with ofatumumab in CLL (PCYC-1109-CA). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 7, 2010—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01217749 NLM Identifier: NCT01217749.

Pharmacyclics, Inc. Ibrutinib and Lenalidomide With Dose Adjusted EPOCH-R in Subjects With Relapsed/Refractory Diffuse Large B-cell Lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). May 12, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02142049?term=NCT02142049 NLM Identifier: NCT02142049.

Pharmacyclics, Inc. Ibrutinib in combination with lenalidomide, with and without rituximab in participants with relapsed or refractory diffuse large B-cell lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Feb. 10, 2014—[cited Apr. 15, 2014]. Available from: http://clinicaltrials.gov/ct2/show/NCT02077166 NLM Identifier: NCT02077166.

Pharmacyclics, Inc. Ibrutinib With Rituximab in Previously Treated Adults With Waldenstrom's Macroglobulinemia. In: ClinicalTrials. gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jun. 9, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02165397?term=NCT02165397 NLM Identifier: NCT02165397.

Pharmacyclics, Inc. Safety and efficacy of PCI-32765 in subjects with relapsed/refractory mantel cell lymphoma (MCL). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 18, 2010—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01236391 NLM Identifier: NCT01236391.

Pharmacyclics, Inc. Safety and efficacy study of Bruton's tyrosine kinase inhibitor in subjects with relapsed or refractory diffuse large B-cell lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Feb. 2, 2011—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01325701 NLM Identifier: NCT01325701.

Pharmacyclics, Inc. Safety and tolerability study of PCI-32765 combined with fludarabine/cyclophosphamide/rituximab (FCR) and bendamustine/rituximab (BR) in chronic lymphocytic leukemia (CLL). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National

#### OTHER PUBLICATIONS

Library of Medicine (US). Feb. 2, 2011—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01292135 NLM Identifier: NCT01292135.

Pharmacyclics, Inc. Safety and tolerability study of PCI-32765 in B Cell lymphoma and chronic lymphocytic leukemia. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Apr. 19, 2010—[cited Nov. 25, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01109069 NLM Identifier: NCT01109069.

Pharmacyclics, Inc. Safety of PCI-32765 in chronic lymphocytic leukemia. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Apr. 13, 2010—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01105247 NLM Identifier: NCT01105247.

Pharmacyclics, Inc. Study of the Bruton's tyrosine kinase inhibitor in combination with carfilzomib (Kyprolis), in subjects with relapsed or relapsed and refractory multiple myeloma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Sep. 27, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01962792 NLM Identifier: NCT01962792.

Pharmacyclics, Inc. Study of the Bruton's tyrosine kinase inhibitor in combination with rituximab in previously untreated subjects with follicular lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 24, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01980654 NLM Identifier: NCT01980654.

Pharmacyclics, Inc. Study of the Bruton's Tyrosine Kinase Inhibitor in Subjects With Chronic Graft Versus Host Disease. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jul. 11, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02195869?term=NCT02195869 NLM Identifier: NCT02195869.

Pharmacyclics, Inc. Study of the Bruton's tyrosine kinase inhibitor in subjects with relapsed or relapsed and refractory multiple myeloma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Nov. 18, 2011—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01478581 NLM Identifier: NCT01478581.

Pharmacyclics, Inc. Study of the Bruton's tyrosine kinase inhibitor in subjects with relapsed/refractory marginal zone lymphoma. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 29, 2013—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT01980628 NLM Identifier: NCT01980628.

Pharmacyclics, Inc. Study of the safety and tolerability of PCI-32765 in patients with recurrent B cell lymphoma (PCYC-04753). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Feb. 20, 2009—[cited Nov. 22, 2013]. Available from: http://clinicaltrials.gov/ct2/show/NCT00849654 NLM Identifier: NCT00849654.

Pharmacyclics: Pharmacyclics initiates phase 1 clinical trial of novel oral Btk inhibitor for refractory B-cell non-Hodgkin's lymphoma. The American Association of Cancer Research (AACR) 100th Annual Meeting in Denver, CO (Apr. 13, 2009).

Pharmacyclics: Study of the Bruton's Tyrosine Kinase Inhibitor in Subjects With Chronic Graft Versus Host Disease. ClinicalTrials. gov [Internet]. Bethesda (MD): National Library of Medicine (US). Available from: http://clinicaltrials.gov/show/NCT02195869.

Picci, P. et al., "Osteosarcoma (osteogenic sarcoma)," Orphanet J Rare Dis. 2(6):1-4 (2007).

Pilchik, "Pharmaceutical Blister Packaging, Part 1: Rationale and Materials," Pharmaceutical Technology, 68-77 (Nov. 2000).

Pileri et al. Mantle Cell Lymphoma. Haematologica 94(11):1488-1492 (2009).

Pollyea et al., "A phase I dose escalation study of the Btk inhibitor PCI-32765 in relapsed and refractory B cell non-Hodgkin lymphoma and use of a novel fluorescent probe pharmacodynamic assay," 51st ASH Annual Meeting and Exposition, Poster Abstract #3713 (2009).

Pollyea et al., "A Phase I Dose Escalation Study of the Btk Inhibitor PCI-32765 in Relapsed and Refractory B Cell Non-Hodgkin Lymphoma and Use of a Novel Fluorescent Probe Pharmacodynamics Assay." Blood. 114(22): Abstract No. 3713 (Nov. 20, 2009).

Ponader et al. The Bruton tyrosine kinase inhibitor PCI-32765 thwarts chronic lymphocytic leukemia cell survival and tissue homing in vitro and in vivo. Blood (Epub Dec. 16, 2011), 119(5):1182-1189 (Feb. 2012).

Powers et al. Irreversible Inhibitors of Serine, Cysteine, and Threonine Proteases. Chem. Rev., 102(12):4639-4750 (2002).

Prakash et al. Chicken sarcoma to human cancers: a lesson in molecular therapeutics. The Ochsner Journal, 7(2):61-64 (Jan. 1, 2007).

Prenata et al., "Separation on the basis of size: Gel permeation chromatography," Protein Purification Methods: A Practical Approach, (Harris & Angal Eds.) IRL Press 1989 293-306.

PRNewsire "U.S. FDA grants regular (full) approval for IMBRUVICA for two indications," Jul. 28, 2014.

PRNewswire, "Pharmacyclics, Inc. announces presentation of interim results from phase I trial of its first-in-human Btk inhibitor PCI-32765," Dec. 7, 2009.

PRNewswire, "Update on preclinical finding and development timeline for PCI-45292," Mar. 2, 2011.

Quek et al. A role for Bruton's tyrosine kinase (Btk) in platelet activation by collagen. Curr. Biol. 8(20):1137-1140 (1998).

Rabin et al. Absolute Lymphocyte Counts Refine MRD-Based Risk Stratification in Pediatric ALL. Blood (Ash Annual Meeting Abstracts) 114:Abstract 1593 (2009).

Rao et al., "Inhibition of Invasion, Angiogenesis, Tumor Growth, and Metastasis by Adenovirus-mediated Transfer of Antisense uPAR and MMP-9 in Non-small Cell Lung Cancer Cells," Mol Cancer Ther, 4(9): 1399-1408 (2005).

Rastetter et al. Rituximab: expanding role in therapy for lymphomas and autoimmune diseases. Ann. Rev. Med 55:477-503 (2004).

Raw et al., "Regulatory Considerations of Pharmaceutical Solid Polymorphism in Abbreviated New Drug Applications (ANDAS)," Adv Drug Deliv Rev, 56: 397-414 (2004).

Remington's Pharmaceutical Sciences, 1995, Mark Printing Co., 17th ed., p. 185.

Ritter et al. Osteosarcoma. Ann. Oncol. 21(Supplement 7):320-325 (2010)

Robak et al. A Targeted Therapy for Protein and Lipid Kinases in Chronic Lymphocytic Leukemia. Curr. Med. Chem. (Epub Jul. 24, 2012), 19(31):5294-5318 (2012).

Robak et al. Tyrosine kinase inhibitors as potential drugs for B-cell lymphoid malignancies and autoimmune disorders. Expert Opin. Investig. Drugs (Epub May 22, 2012), 21(7):921-947 (Jul. 2012). Rushworth et al. BTK inhibitor ibrutinib is cytotoxic to myeloma and potently enhances bortezomib and lenalidomide activities through NF-κb. Cell Signal (Epub Sep. 11, 2012), 25(1):106-112 (Jan. 2013).

Sada et al. Protein-Tyrosine Kinases and Adaptor Proteins in FceRI-Mediated Signaling in Mast Cells. Curr. Mol. Med. 3(1):85-94 (2003).

Saulnier et al., "An efficient method for the synthesis of guanidine prodrugs," Bioorganic and Medicinal Chemistry Letters, vol. 4, p. 1985-1990 (1994).

Schaeffer et al. Tec family kinases in lymphocyte signaling and function. Curr. Op. Imm. 12:282-288 (2000).

Schnute et al., "Bruton's tyrosine kinase (Btk)," Anti-Inflammatory Drug Discovery, Ed. J.I. Levin and S. Laufer, (2012), pp. 297-326. Schwamb et al. B-cell receptor triggers drug sensitivity of primary CLL cells by controlling glucosylation of ceramides. Blood (Epub Aug. 27, 2012), 120(19):3978-3985 (Nov. 2012).

Science Daily Counting tumor cells in blood predicts treatment benefit in prostate cancer. (Jul. 7, 2008), http://www.sciencedaily.com/releases/2008/07/080706083142.htm , last accessed Jul. 23, 2013.

#### OTHER PUBLICATIONS

Science Daily, "Drug shows surprising efficacy as treatment for chronic leukemia, mantle cell lymphoma," (Jun. 19, 2013), http://www.sciencedaily.com/releases/2013/06/130619195217.htm, last accessed Jan. 15, 2016.

Science IP CAS Search, Mar. 16, 2006.

Science IP CAS Search, Sep. 5, 2006.

Shaffer et al. Lymphoid malignancies: the dark side of B-cell differentiation. Nature Reviews/Immunology 2:920-932 (2002).

Shah et al. Ibrutinib for the treatment of mantle cell lymphoma. Expert Rev. Hematol. 7(5):521-531 (2014) (Epub Aug. 27, 2014). Silverman. The Organic Chemistry of Drug Design and Drug Action, Academic Press, Inc., San Diego, pp. 352-401 (1992).

Sinkula et al. Rationale for design of biologically reversible drug derivatives: prodrugs. J. Pharm. Sci., 64:181-210 (1975).

Sivina et al. CCL3 (MIP-1a) Plasma Levels and The Risk for Disease Progression in Chronic Lymphocytic Leukemia. Blood 117(5):1662-1669 (2010).

Smaill et al., "Irreversible Inhibitors of the Epidermal Growth Factor Receptor: 4-(Phenylamino)quinazoline- and 4-(Phenylamino)pyrido[3,2-d]pyrimidine-6-acrylamides Bearing Additional Solubilizing Functions," J Med Chem, 43: 1380-1397 (2000).

Smaill et al., "Tyrosine kinase inhibitors. 15. 4-(phenylamino)quinazoline and 4-(phenylamino)pyrido[d]pyrimidine acrylamides as irreversible inhibitors of the ATP binding site of the epidermal growth factor receptor," J Med Chem, 42(10):1803-15 (1999).

Smith et al., "The Tec family of cytoplasmic tyrosine kinases: mammalian Btk, Bmx, Itk, Tec, Txk and homologs in other species," BioEssays, 23:436-46 (2001).

Smolen et al. Therapeutic Strategies for Rheumatoid Arthritis. Nature Reviews 2:473-488 (2003).

Stead et al. Concise synthesis of (+/-)-Cytisine via lithiation of N-Boc-bispidine. Organic Letters 7(20):4459-4462 (2005).

STN Registry No. 936563-96-1. Ibrutinib. Retrieved from STN Registry Jul. 27, 2015. 1 pg.

Strimbu et al. What are biomarkers? Curr Opin HIV AIDS 5(6):463-466 (2010.

Supplementary European Search Report EP13799982.7 dated Dec. 14, 2015.

Supplementary European Search Report for EP13854424 dated Mar. 14, 2016.

Taiwan Search Report for Application No. 102119583, dated Oct. 24, 2016.

Takahashi et at. Serum CCL3 and CCL4 Levels Function as Novel Prognostic Markers in Diffuse Large B Cell Lymphoma [online]. 54th ASH Annual Meeting and Exposition. [retrieved on Apr. 21, 2015], Abstract 2709. Retrieved from the Internet. URL: https://ash.confex.com/ash/2012/webprogram/Paper53900.html

TG Therapeutics, Inc. Ublituximab + ibrutinib in select B-cell malignancies. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Dec. 11, 2013—[cited Apr. 15, 2014]. Available from: http://clinicaltrials.gov/ct2/show/NCT02013128 NLM Identifier: NCT02013128.

The Lymphoma Academic Research Organisation. Bruton's tyrosine kinase (BTK) inhibition in B-cell lymphomas (BIBLOS). In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Jan. 31, 2014—[cited Apr. 15, 2014]. Available from: http://clinicaltrials.gov/th/show/NCT02055924 NLM Identifier: NCT02055924

Tinmouth et al. Fludarabine in alkylator-resistant follicular non-Hodgkin's lymphoma. Leuk. Lymphoma 41(1-2):137-145 (2001). Traxler et al., "Use of a pharmacophore model for the design of EGF-R tyrosine kinase inhibitors: 4-(phenylamino)pyrazolo[3,4-d]pyramidines," J Med Chem, 40(22):3601-16 (1997).

Uckun et al. Bruton's Tyrosine Kinase (BTK) as a Dual-Function Regulator of Apoptosis. Biochem. Pharmacology 56:683-691 (1998).

Uckun et al. The Anti-leukemic Bruton's Tyrosin Kinase Inhibitor a-cyano-β-hydroxy-β-mehyl-N-(2,5-dibromophenyl)Propenamide (LFM-A13)Prevents Fatal Thromboembolism. Leuk. Lymphoma 44(9):1569-1577 (2003).

Uckun et al., "Bruton's tyrosine kinase as a molecular target in treatment of leukemias and lymphomas as well as inflammatory disorders and autoimmunity," Expert Opin Ther Pat, 20(11):1-14 (2010).

Uckun et al., "Btk as a mediator of radiation-induced apoptosis in DT-40 lymphoma B cells," Science, 273(5278):1096-100 (1996). Uckun et al., "In vivo pharmacokinetic features, toxicity profile, and chemosensitizing activity of alpha-cyano-beta-hydroxy-beta-methyl-N-(2,5-dibromophenyl)propenamide (LFM-A13), a novel antileukemic agent targeting Bruton's tyrosine kinase," Clin Cancer Res, 8(5):1224-33 (2002).

University of California, San Diego. A Phase lb/II Study of Ibrutinib in Combination With GA101—Obinutuzumab in Previously Untreated Chronic Lymphocytic Leukemia (CLL) Patients Over 65 Years of Age or With Comorbidities That Preclude the Use of Chemotherapy Based Treatment. In: ClinicalTrials.gov [Internet]. Bethesda (MD): National Library of Medicine (US). Oct. 30, 2014 [cited Feb. 5, 2015] Available from: https://clinicaltrial.gov/ct2/show/NCT02315768?term=NCT02315768 NLM Identifier: NCT02315768

Vassilev et al. Bruton's Tyrosine Kinase as an Inhibitor of the Fas/CD95 Death-inducing Signaling Complex. J. Biol. Chem. 274(3):1646-1656 (1999).

Vassilev et al. Therapeutic Potential of Inhibiting Bruton's Tyrosine Kinase, (BTK). Current Pharmaceutical Design 10:1757-1766 (2004).

Vippagunta et al. Crystalline Solids, Advanced Drug Delivery Reviews. 48:3-26 (2001).

Vose. Mantle cell lymphoma: 2012 update on diagnosis, risk-stratification, and clinical management. Am. J. Hematol. 87(6):604-609 (Jun. 2012).

Wang et al. "Ibrutinib and rituximab are an efficacious and safe combination in relapsed mantle cell lymphoma: preliminary results from a Phase II clinical trial," Oral Abstract Session 624, 56th ASH Annual Meeting and Exposition (Dec. 6-9, 2014).

Wang et al. Targeting BTK with ibrutinib in relapsed or refractory mantel-cell lymphoma. N Engl J Med 369(6):507-516 (Aug. 8, 2013).

Wilkinson et al. Selective tyrosine kinase inhibitors. Expert Opin. Emerging Drugs 5(3):287-297 (2000).

Wilson et al., "The Bruton's tyrosine kinase (BTK) inhibitor, ibrutinib (PCI-32765), has preferential activity in the ABC subtype of relapsed/refractory de novo diffuse large B-cell lymphoma (DLBCL): interim results of a multicenter, open-label, phase 2 study," Blood 120:Abstract 686 (2012).

Witzens-Harig et al. Current treatment of mantle cell lymphoma: results of a national survey and consensus meeting. Ann Hematol. (Epub Aug. 29, 2012), 91(11):1765-1772 (Nov. 2012).

Witzig et al. Lenalidomide oral monotherapy produces durable responses in relapsed or refractory indolent non-Hodgkin's lymphoma. J. Clin. Oncol. 27:5404-5409 (Epub Oct. 5, 2009).

Wolff. Burger's Medicinal Chemistry and Drug Discovery. 5th Ed. Part 1, pp. 975-977 (1995).

Woyach et al., "Resistance Mechanisms for the Bruton's Tyrosine Kinase Inhibitor Ibrutinib," N Engl J Med, 370(24): 2286-2294 (2014).

Yamamoto et al. The Orally Available Spleen Tyrosine Kinase Inhibitor 2-[7-(3,4-Dimethoxyphenyl)-imidazo[1,2-c]pyrimidin-5-ylamino]-nicotinamide Dihydrochloride (BAY61-3606) Blocks Antigen-Induced Airway Inflammation in Rodents. J. Pharma. and Exp. Therapeutics 306(3):1174-1181 (2003).

Yang et al. Tyrosine kinase inhibition in diffuse large B-cell lymphoma: molecular basis for antitumor activity and drug resistance of dasatinib. Leukemia 22(9):1755-1766 (2008) [E-pub Jul. 3, 2008]. Yasuhiro et al., "ONO-WG-307, a novel, potent and selective inhibitor of Bruton's tyrosine kinase, in sustained inhibition of the Erk, Akt and PKD signaling pathways," 53rd American Society of Hematology Annual Meeting and Exposition, San Diego, CA, Poster #2021 (2011).

#### OTHER PUBLICATIONS

Zent et al. The Treatment of Recurrent/Refractory chronic Lymphocytic Leukemia/Small Lymphocytic Lymphoma (CLL) With Everolimus Results in Clinical Responses and Mobilization of CLL Cells Into the Circulation. Cancer 116(9):2201-2207 (2010). Zhu et al., "Calpain inhibitor II induces caspase-dependent apoptosis in human acute lymphoblastic leukemia and non-Hodgkin's lymphoma cells as well as some solid tumor cells," Clin Cancer Res, 6:2456-63 (2000)

Zhu T., et al., "New Tablet Formulation of Lopinavir/Ritonavir is Bioequivalent to the Capsule at a Dose of 800/200 mg," Poster H-1894, 45th Interscience Conference on Antimicrobial Agents and Chemotherapy (ICAAC), Dec. 16-19, 2005, 4 pages. Zvonicek et al., "First Crystal Structures of Pharmaceutical

Ibrutinib: Systematic Solvate Screening and Characterization," Crystal Grwoth & Design, 17(6): 3116-3127 (2017). Ahead of print. "Imbruvica," EPAR Summary for the Public, European Medicines Agency Science Medicines Health last updated Aug. 2016, retrieved from: <a href="mailto://www.ema.europa.eu/ema/index.jsp?curl=pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages/medi-pages cines/human/medicines/003791/human\_med\_001801.jsp

&mid=WC0b01ac058001d124>.

Label for IMBRUVICA® (ibrutinib), revised Feb. 2018.

Supplementary European Search Report for EP 15829513.9, dated Mar. 7, 2018.

Teja et al., "Drug-excipient behavior in polymeric amorphous solid dispersions," Journal of Excipients and Food Chemicals, 4(3): 70-94 (2013).

Vasconcelos et al., "Solid dispersions as strategy to improve oral bioavailability of poor water soluble drugs," Drug Discov Today, 12(23-24):1068-1075 (2007).

<sup>\*</sup> cited by examiner

FIG. 1 30 Mean Ibrutinib Plasma Concentration (ng/ml.) Formulation A Formulation 8 25 Formulation C Formulation D 20 15 10 12 6 18 24 Nominal Time Post-Dose (h)

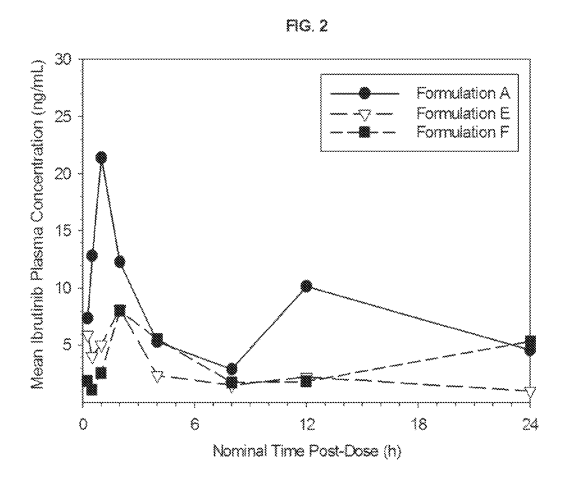
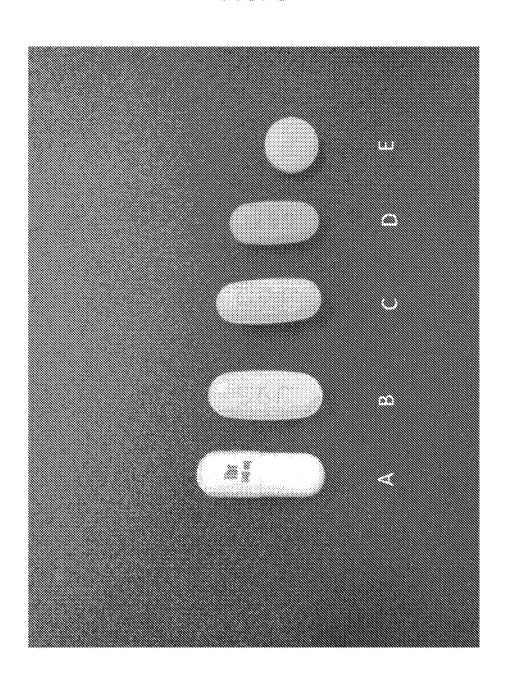


FIG. 3



# PHARMACEUTICAL FORMULATIONS OF A BRUTON'S TYROSINE KINASE INHIBITOR

## CROSS-REFERENCE TO RELATED APPLICATION

This application is a continuation of U.S. patent application Ser. No. 15/862,995, filed Jan. 5, 2018, which is a continuation of U.S. patent application Ser. No. 15/467,414, filed Mar. 23, 2017, now abandoned, which is a continuation of U.S. patent application Ser. No. 15/060,010, filed Mar. 3, 2016, now U.S. Pat. No. 9,655,857, issued May 23, 2017, which claims the benefit of U.S. Provisional Application No. 62/127,717, filed Mar. 3, 2015, and U.S. Provisional Application No. 62/193,518, filed Jul. 16, 2015, which are each incorporated herein by reference in their entireties.

#### FIELD OF THE INVENTION

Described herein is the Bruton's tyrosine kinase (Btk) <sup>20</sup> inhibitor 1-((R)-3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl)piperidin-1-yl)prop-2-en-1-one, including pharmaceutical compositions, solvates and pharmaceutical formulations that include the Btk inhibitor and methods of using the Btk inhibitor compositions or formulations in the treatment of diseases or conditions that would benefit from inhibition of Btk activity.

#### BACKGROUND OF THE INVENTION

Bruton's tyrosine kinase (Btk), a member of the Tec family of non-receptor tyrosine kinases, is a key signaling enzyme expressed in all hematopoietic cells types except T lymphocytes and natural killer cells. Btk plays an essential <sup>35</sup> role in the B-cell signaling pathway linking cell surface B-cell receptor (BCR) stimulation to downstream intracellular responses.

Btk is a key regulator of B-cell development, activation, signaling, and survival. In addition, Btk plays a role in a 40 number of other hematopoietic cell signaling pathways, e.g., Toll like receptor (TLR) and cytokine receptor-mediated TNF-α production in macrophages, IgE receptor (FcepsilonRI) signaling in Mast cells, inhibition of Fas/APO-1 apoptotic signaling in B-lineage lymphoid cells, and collagen-stimulated platelet aggregation.

1-((R)-3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3, 4-d]pyrimidin-1-yl)piperidin-1-yl)prop-2-en-1-one is also known by its IUPAC name as 1-{(3R)-3-[4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl]piperidin-1-yl}prop-2-en-1-one or 2-Propen-1-one, 1-[(3R)-3-[4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl]-1-piperidinyl-, and has been given the USAN name, ibrutinib. The various names given for ibrutinib are used interchangeably herein.

#### SUMMARY OF THE INVENTION

Described herein is the Btk inhibitor 1-((R)-3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl)pi-600 peridin-1-yl)prop-2-en-1-one, including pharmaceutically acceptable compositions, formulations, and methods of uses thereof. Also described are pharmaceutically acceptable compositions and formulations of the Btk inhibitor, 1-((R)-3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]py-7 rimidin-1-yl)piperidin-1-yl)prop-2-en-1-one, used in the manufacture of medicaments for the treatment of diseases or

2

conditions that are associated with Btk activity. 1-((R)-3-(4-Amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl)piperidin-1-yl)prop-2-en-1-one is an irreversible Btk inhibitor. Further described are pharmaceutical compositions and formulations of the Btk inhibitor, 1-((R)-3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl)piperidin-1-yl)prop-2-en-1-one, and methods of using the Btk inhibitor in the treatment of diseases or conditions (including diseases or conditions wherein irreversible inhibition of Btk provides therapeutic benefit to a mammal having the disease or condition).

Also described herein is a process for preparing a pharmaceutical composition of 1-((R)-3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl)piperidin-1-yl)prop-2-en-1-one by a wet granulation method. Further described are pharmaceutical formulations that include a pharmaceutical composition of 1-((R)-3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl)piperidin-1-yl)prop-2-en-1-one prepared by a wet granulation method.

In one aspect is a pharmaceutical composition comprising ibrutinib, wherein ibrutinib is a compound with the structure of Compound 1,

Compound 1  $NH_2$  ,

and wherein the pharmaceutical composition comprises at least 50% w/w of ibrutinib.

In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises about 50% w/w to about 90% w/w of ibrutinib. In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises about 50% w/w to about 80% w/w of ibrutinib. In another embodiment is a pharmaceutical com-55 position comprising ibrutinib, wherein the pharmaceutical composition comprises about 60% w/w to about 80% w/w of ibrutinib. In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises about 60% w/w to about 75% w/w of ibrutinib. In another embodiment is a pharmaceutical composition comprising at least 50% w/w of ibrutinib, wherein the pharmaceutical composition comprises intragranular and extragranular ingredients. In another embodiment is a pharmaceutical composition comprising at least 50% w/w of ibrutinib, wherein the pharmaceutical composition is prepared using a wet granulation method. In another embodiment is a pharmaceutical composition comprising at least

50% w/w of ibrutinib, further comprising at least one pharmaceutically acceptable excipient.

In another embodiment is a high-load solid tablet formulation comprising a pharmaceutical composition comprising at least 50% w/w of ibrutinib, about 50% w/w to about 90% 5 w/w of ibrutinib, about 50% w/w to about 80% w/w of ibrutinib, about 60% w/w to about 80% w/w of ibrutinib, or about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a high-load solid tablet formulation compris- 10 ing a pharmaceutical composition comprising at least 50% w/w of ibrutinib, about 50% w/w to about 90% w/w of ibrutinib, about 50% w/w to about 80% w/w of ibrutinib, about 60% w/w to about 80% w/w of ibrutinib, or about 60% w/w to about 75% w/w of ibrutinib, and one or more 15 pharmaceutically acceptable excipients wherein the one or more excipients are present in an amount from about 10% w/w to about 50% w/w.

In another embodiment is a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, about 50% 20 w/w to about 90% w/w of ibrutinib, about 50% w/w to about 80% w/w of ibrutinib, about 60% w/w to about 80% w/w of ibrutinib, or about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are selected from the 25 group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants. In some embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, mannitol, xylitol, 30 sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is present in an amount from 35 about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrystalline cellulose. In 40 some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 45 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent 50 comprises lactose and microcrystalline cellulose. In some embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an 55 amount of about 14% w/w and microcrystalline cellulose is present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, 60 a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose, croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymethylcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, 65 cross-linked polymer such as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In

4

some embodiments, the disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to about 10% w/w, about 5% w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the polyvinylpyrrolidone is present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipient is a surfactant. In some embodiments, the surfactant is sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some embodiments, at least one excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium stearate is present in an amount from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise, consist essentially of, or consist, lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide and magnesium stearate. In another embodiment, the excipients comprise, consist essentially of, or consist, lactose, polyvinylpyrrolidone, sodium lauryl sulfate, crospovidone, colloidal silicon dioxide, and magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, about 50% w/w to about 90% w/w of ibrutinib, about 50% w/w to about 80% w/w of ibrutinib, about 60% w/w to about 80% w/w of ibrutinib, or about 60% w/w to about 75% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise, consist essentially of, or consist lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment is a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, about 50% w/w to about 90% w/w of ibrutinib, about 50% w/w to about 80% w/w of ibrutinib, about 60% w/w to about 80% w/w of ibrutinib, or about 60% w/w to about 75% w/w of ibrutinib, wherein the intragranular excipients comprise

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;

- microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w;
- croscarmellose sodium in an amount from about 0 to about 10% w/w, about 2% w/w to about 5% w/w, or 5 about 2% w/w to about 4% w/w; and
- hydroxypropyl cellulose in an amount from about 0% w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and

#### the extragranular excipients comprise

- croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 5% w/w;
- sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w;
- colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and
- magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment is a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, about 50% 25 w/w to about 90% w/w of ibrutinib, about 50% w/w to about 80% w/w of ibrutinib, about 60% w/w to about 80% w/w of ibrutinib, or about 60% w/w to about 75% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose, microcrystalline 30 cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment is a high-load solid tablet formulation compris- 35 ing at least 50% w/w of ibrutinib, about 50% w/w to about 90% w/w of ibrutinib, about 50% w/w to about 80% w/w of ibrutinib, about 60% w/w to about 80% w/w of ibrutinib, or about 60% w/w to about 75% w/w of ibrutinib, wherein the intragranular excipients comprise

- lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;
- microcrystalline cellulose in an amount from about 1% w/w to about 10% w/w, about 2% w/w to about 5% w/w;
- polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;
- croscarmellose sodium in an amount from about 1% w/w to about 10% w/w, or about 3% w/w to about 7% w/w; and
- sodium lauryl sulfate in an amount from about 0 to about 2% w/w, about 0.5% w/w to about 1.5% w/w; and the extragranular excipients comprise
  - croscarmellose sodium in an amount from about 0 to about 5% w/w, about 1% w/w to about 3% w/w;
  - sodium lauryl sulfate in an amount from about 0 to about 10% w/w or about 0% w/w to about 4% w/w;
  - colloidal silicon dioxide in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and
  - magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment is a high-load solid tablet formulation comprising

- a) about 69% w/w to about 71% w/w of ibrutinib,
- b) about 13% w/w to about 15% w/w of lactose,

6

- c) about 2% w/w to about 5% w/w of microcrystalline cellulose.
- d) about 1% w/w to about 3% w/w of polyvinylpyrrolidone.
- e) about 6% w/w to about 8% w/w of croscarmellose sodium.
- f) about 1% w/w to about 4% w/w of sodium lauryl sulfate,
- g) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
- h) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising

- a) about 70% w/w of ibrutinib,
- b) about 14% w/w of lactose monohydrate,
- c) about 5% w/w of microcrystalline cellulose,
- d) about 2% w/w of polyvinylpyrrolidone,
- e) about 7% w/w of croscarmellose sodium,
- f) about 1% w/w of sodium lauryl sulfate,
- g) about 0.5% w/w of colloidal silicon dioxide, and
- h) about 0.5% w/w of magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising

- a) about 70% w/w of ibrutinib,
- b) about 14% w/w of lactose monohydrate,
- c) about 2% w/w of microcrystalline cellulose,
- d) about 2% w/w of polyvinylpyrrolidone,
- e) about 7% w/w of croscarmellose sodium,
- f) about 4% w/w of sodium lauryl sulfate,
- g) about 0.5% w/w of colloidal silicon dioxide, and
- h) about 0.5% w/w of magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising

- a) about 70% w/w of ibrutinib,
- b) about 16% w/w of lactose,
- c) about 2% w/w of polyvinylpyrrolidone,
- d) about 1% w/w of sodium lauryl sulfate,
- e) about 10% w/w of crospovidone,
- f) about 0.5% w/w of colloidal silicon dioxide, and
- g) about 0.5% w/w of magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising

- a) about 59% w/w to about 61% w/w of ibrutinib,
- b) about 13% w/w to about 15% w/w of lactose,
- about 13% w/w to about 15% w/w of microcrystalline cellulose,
- d) about 4% w/w to about 6% w/w of croscarmellose sodium,
- e) about 5% w/w to about 7% w/w of sodium lauryl sulfate.
- f) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
- g) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In some embodiments, the total weight of a tablet is about 934 mg.

In another embodiment is a high-load solid tablet formulation comprising

- a) about 59% w/w to about 61% w/w of ibrutinib,
- b) about 13% w/w to about 14% w/w of lactose,

65

- c) about 13% w/w to about 14% w/w of microcrystalline cellulose.
- d) about 2% w/w to about 3% w/w of croscarmellose sodium (intragranular),
- e) about 0.8% w/w to about 1.2% w/w of hydroxypropyl cellulose,

- f) about 2% w/w to about 3% w/w of croscarmellose sodium (extragranular),
- g) about 5.5 to about 6.5% w/w of sodium lauryl sulfate,
- h) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
- i) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In some embodiments, the total weight of a tablet is about 934 mg.

In another embodiment is a high-load solid tablet formulation comprising

- a) about 69% w/w to about 71% w/w of ibrutinib,
- b) about 8% w/w to about 9% w/w of lactose,
- c) about 8 to about 9% w/w of microcrystalline cellulose,
- d) about 2.5 to about 3.5% w/w of croscarmellose sodium 15 (intragranular),
- e) about 2.5 to about 3.5% w/w of croscarmellose sodium (extragranular),
- g) about 5.5 to about 6.5% w/w of sodium lauryl sulfate,
- h) about 0.4% w/w to about 0.6% w/w of colloidal silicon 20 dioxide, and
- i) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

Lactose used herein may be anhydrous lactose and/or hydrous lactose, such as lactose monohydrate. In some 25 embodiments, the lactose is anhydrous lactose. In some particular embodiments, the lacose is hydrous lactose. In more particular embodiments, the lacose is lactose monohydrate.

In some embodiments, the total weight of a tablet is about 30 800 mg.

In some embodiments of the high-load solid tablet formulations described herein, the ibrutinib is in an amount of about 35 mg to about 840 mg per tablet. In some embodiments of the high-load solid tablet formulations described 35 herein, the ibrutinib is in an amount of about 140 mg to about 840 mg per tablet. In some embodiments of the high-load solid tablet formulations described herein, the ibrutinib is in an amount of about 140 mg, about 280 mg, about 420 mg, about 560 mg, or about 840 mg per tablet, or 40 any range between any two of the values, end points inclusive. In some embodiments of the high-load solid tablet formulations described herein, the ibrutinib is in an amount of about 560 mg. In some embodiments of the high-load solid tablet formulations described herein, the ibrutinib is in 45 micronized form. In some embodiments of the high-load solid tablet formulations described herein, the formulation is used for once a day dosing. In some embodiments of the high-load solid tablet formulations described herein, the formulation is in an oral dosage form containing a thera- 50 peutically effective amount of ibrutinib.

In another embodiment is a method of treating a disease in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described 55 herein.

In another embodiment is a method of treating an autoimmune disease in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein. In some embodiments, the autoimmune disease is rheumatoid arthritis or lupus.

In another embodiment is a method of treating a heteroimmune disease in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

8

In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein. In some embodiments, the cancer is a B-cell proliferative disorder. In some embodiments, the cancer is a B-cell proliferative disorder and the B-cell proliferative disorder is diffuse large B cell lymphoma, follicular lymphoma or chronic lymphocytic leukemia. In some embodiments, the cancer is a B cell malignancy. In some embodiments, the cancer is a B cell malignancy and the B cell malignancy selected from chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL), mantle cell lymphoma (MCL), diffuse large B Cell lymphoma (DLBCL), and multiple myeloma. In some embodiments, the cancer is a lymphoma, leukemia or a solid tumor. In some embodiments, the cancer is diffuse large B cell lymphoma, follicular lymphoma, chronic lymphocytic lymphoma, chronic lymphocytic leukemia, B-cell prolymphocytic leukemia, lymphoplasmacytic lymphoma/Waldenstrom macroglobulinemia, splenic marginal zone lymphoma, plasma cell myeloma, plasmacytoma, extranodal marginal zone B cell lymphoma, nodal marginal zone B cell lymphoma, mantle cell lymphoma, mediastinal (thymic) large B cell lymphoma, intravascular large B cell lymphoma, primary effusion lymphoma, burkitt lymphoma/leukemia, or lymphomatoid granulomatosis.

In another embodiment is a method of treating mastocytosis in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

In another embodiment is a method of treating osteoporosis or bone resorption disorders in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

In another embodiment is a method of treating an inflammatory disease or condition in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

In another embodiment is a method of treating lupus in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

In some embodiments, the formulations and methods described herein can be used to treat carcinoma of the brain, kidney, liver, adrenal gland, bladder, breast, stomach, gastric tumors, ovaries, colon, rectum, prostate, pancreas, lung, vagina, cervix, testis, genitourinary tract, esophagus, larynx, skin, bone or thyroid, sarcoma, glioblastomas, neuroblastomas, multiple myeloma, gastrointestinal cancer, especially colon carcinoma or colorectal adenoma, a tumor of the neck and head, an epidermal hyperproliferation, psoriasis, prostate hyperplasia, a neoplasia, a neoplasia of epithelial character, adenoma, adenocarcinoma, keratoacanthoma, epidermoid carcinoma, large cell carcinoma, non-small-cell lung carcinoma, lymphomas, Hodgkins and Non-Hodgkins, a mammary carcinoma, follicular carcinoma, undifferentiated carcinoma, papillary carcinoma, seminoma, melanoma, or Smoldering of indolent multiple myeloma.

In some embodiments, the formulations and methods described herein can be used to treat a central nervous system (CNS) malignancy. In some embodiments, the CNS malignancy is a primary CNS lymphoma. In some embodiments the primary CNS lymphoma is a glioma. In some

embodiments the glioma is astrocytomas, ependymomas, oligodendrogliomas. In some embodiments the CNS malignancy is astrocytic tumors such as juvenile pilocytic, subependymal, well differentiated or moderately differentiated anaplastic astrocytoma; anaplastic astrocytoma; glioblastoma multiforme; ependymal tumors such as myxopapillary and well-differentiated ependymoma, anaplastic ependymoma, ependymoblastoma; oligodendroglial tumors including well-differentiated oligodendroglioma and anaplastic oligodendroglioma; mixed tumors such as mixed astrocytoma-ependymoma, mixed astrocytoma-oligodendroglioma, mixed astrocytoma-oligodendroglioma; or medulloblastoma.

In some embodiments, the formulations and methods described herein can be used to treat hematological malignancies such as, but not limited to, a leukemia, a lymphoma, a myeloma, a non-Hodgkin's lymphoma, a Hodgkin's lymphoma, a T-cell malignancy, or a B-cell malignancy. In some embodiments, the hematological malignancy is a treatment païve hematological malignancy. In some embodiments the hematological malignancy is a relapsed or refractory hematological malignancy.

In some embodiments, the hematologic malignancy is a T-cell malignancy. In some embodiments, the T-cell malignancy is peripheral T-cell lymphoma not otherwise specified (PTCL-NOS), anaplastic large cell lymphoma, angioimmunoblastic lymphoma, cutaneous T-cell lymphoma, adult T-cell leukemia/lymphoma (ATLL), blastic NK-cell lymphoma, enteropathy-type T-cell lymphoma, hematosplenic gamma-delta T-cell lymphoma, lymphoblastic lymphoma, nasal NK/T-cell lymphomas, or treatment-related T-cell lymphomas. In some embodiments, the T-cell malignancy is a relapsed or refractory T-cell malignancy. In some embodiments, the T-cell malignancy is a treatment naïve T-cell malignancy.

In some embodiments, the hematologic malignancy is a B-cell proliferative disorder. In some embodiments, the cancer is chronic lymphocytic leukemia (CLL), small lym- 40 phocytic lymphoma (SLL), high risk CLL, a non-CLL/SLL lymphoma, or prolymphocytic leukemia (PLL). In some embodiments, the cancer is follicular lymphoma (FL), diffuse large B-cell lymphoma (DLBCL), mantle cell lymphoma (MCL), Waldenstrom's macroglobulinemia, multiple 45 myeloma, extranodal marginal zone B cell lymphoma, nodal marginal zone B cell lymphoma, Burkitt's lymphoma, non-Burkitt high grade B cell lymphoma, primary mediastinal B-cell lymphoma (PMBL), immunoblastic large cell lymphoma, precursor B-lymphoblastic lymphoma, B cell pro- 50 lymphocytic leukemia, lymphoplasmacytic lymphoma, splenic marginal zone lymphoma, plasma cell myeloma, plasmacytoma, mediastinal (thymic) large B cell lymphoma, intravascular large B cell lymphoma, primary effusion lymphoma, or lymphomatoid granulomatosis. In some embodi- 55 ments, DLBCL is further divided into subtypes: activated B-cell diffuse large B-cell lymphoma (ABC-DLBCL), germinal center diffuse large B-cell lymphoma (GCB DLBCL), and Double-Hit (DH) DLBCL. In some embodiments, ABC-DLBCL is characterized by a CD79B mutation. In some 60 embodiments, ABC-DLBCL is characterized by a CD79A mutation. In some embodiments, the ABC-DLBCL is characterized by a mutation in MyD88, A20, or a combination thereof. In some embodiments, the cancer is acute or chronic myelogenous (or myeloid) leukemia, myelodysplastic syn- 65 drome, or acute lymphoblastic leukemia. In some embodiments, the B-cell proliferative disorder is a relapsed and

10

refractory B-cell proliferative disorder. In some embodiments, the B-cell proliferative disorder is a treatment naïve B-cell proliferative disorder.

In some embodiments, the formulations and methods described herein can be used to treat fibrosis. In some embodiments, the fibrosis is not associated with graft versus host disease (GVHD). In some embodiments, the fibrosis is not associated with sclerodermatous GVHD, lung chronic GVHD, or liver chronic GVHD. In some embodiments, the fibrosis is of the liver, lung, pancreas, kidney, bone marrow, heart, skin, intestine, or joints. In some embodiments, the fibrosis is of the lung. In some embodiments, the fibrosis is of the lung. In some embodiments, the fibrosis is of the pancreas. In some embodiments, the patient has cirrhosis, chronic pancreatitis, or cystic fibrosis.

In another aspect is a process for preparing a pharmaceutical composition or tablet formulation comprising ibrutinib as described herein, wherein the process comprises a wet granulation method.

In another aspect is a high-load solid tablet formulation comprising ibrutinib, wherein ibrutinib is a compound with the structure of Compound 1,

 $NH_2$   $NH_2$  N N N

Compound 1

and the tablet comprises about 560 mg of ibrutinib.

In another embodiment is a high-load solid tablet formulation, wherein ibrutinib is in micronized form. In another embodiment, ibrutinib is in spray-dried form. In another embodiment, ibrutinib is not in spray-dried form. In another embodiment, the particle size is about or less than 30 micron. In one embodiment, ibrutinib is in micronized form and the particle size is about 1-30 micron. In another embodiment, the particle size is about or less than 10 micron. In another embodiment, the particle size is <1 micron. In another embodiment is a high-load solid tablet formulation, wherein the tablet is used for once a day oral dosing.

In another aspect, provided herein are methods for treating a patient by administering Compound 1. In some embodiments, provided herein is a method of inhibiting the activity of tyrsoine kinase(s), such as Btk, or of treating a disease, disorder, or condition, which would benefit from inhibition of tyrosine kinase(s), such as Btk, in a mammal, which includes administering to the mammal a therapeutically effective amount of Compound 1, or pharmaceutically acceptable salt, pharmaceutically active metabolite, pharmaceutically acceptable prodrug, or pharmaceutically acceptable solvate.

In another aspect, provided herein is the use of Compound 1 for inhibiting Bruton's tyrosine kinase (Btk) activity or for the treatment of a disease, disorder, or condition, which would benefit from inhibition of Bruton's tyrosine kinase (Btk) activity.

In some embodiments, a pharmaceutical composition comprising crystalline Compound 1 is administered to a human. In some embodiments, a pharmaceutical composition comprising amorphous Compound 1 is administered to a human.

In some embodiments, a pharmaceutical composition comprising crystalline Compound 1 is orally administered. In some embodiments, a pharmaceutical composition comprising amorphous Compound 1 is orally administered.

In some embodiments, a pharmaceutical composition 15 comprising crystalline Compound 1 is used for the formulation of a medicament for the inhibition of tyrosine kinase activity. In some other embodiments, a pharmaceutical composition comprising crystalline Compound 1 is used for the formulation of a medicament for the inhibition of Bruton's 20 tyrosine kinase (Btk) activity. In some embodiments, a pharmaceutical composition comprising amorphous Compound 1 is used for the formulation of a medicament for the inhibition of tyrosine kinase activity. In some other embodiments, a pharmaceutical composition comprising amor- 25 phous Compound 1 is used for the formulation of a medicament for the inhibition of Bruton's tyrosine kinase (Btk) activity.

In some embodiments, in any of the embodiments disclosed herein (including compositions, methods, uses, formulations, combination therapy, etc.), Compound 1, or a pharmaceutically acceptable salt or solvate thereof, is optically pure (i.e. greater than 99% chiral purity by HPLC). In some embodiments, in any of the embodiments disclosed herein (including compositions, methods, uses, formula- 35 tions, combination therapy, etc.), Compound 1, or a pharmaceutically acceptable salt or solvate thereof, is replaced with: a) Compound 1, or a pharmaceutically acceptable salt or solvate thereof, of lower chiral purity; b) 1-((S)-3-(4-1-yl)piperidin-1-yl)prop-2-en-1-one, or a pharmaceutically acceptable salt or solvate thereof of any optical purity; or c) racemic 1-(3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo [3,4-d]pyrimidin-1-yl)piperidin-1-yl)prop-2-en-1-one, or a pharmaceutically acceptable salt or solvate thereof.

In any of the embodiments disclosed herein (including compositions, methods, uses, formulations, combination therapy, etc.), amorphous Compound 1 is used. In any of the embodiments disclosed herein (including compositions, methods, uses, formulations, combination therapy, etc.), 50 crystalline Compound 1 is used.

In some embodiments, in any of the embodiments disclosed herein (including compositions, methods, uses, formulations, combination therapy, etc.), Compound 1, or a pharmaceutically acceptable salt thereof, is replaced with an 55 active metabolite of Compound 1. In some embodiments, the active metabolite is in a crystalline form. In some embodiments, the active metabolite is in an amorphous phase. In further embodiments the metabolite is isolated. In some embodiments, in any of the embodiments disclosed 60 herein (including compositions, methods, uses, formulations, combination therapy, etc.), Compound 1, or a pharmaceutically acceptable salt thereof, is replaced with a prodrug of Compound 1, or a deuterated analog of Compound 1, or a pharmaceutically acceptable salt thereof.

Other objects, features and advantages of the methods and compositions described herein will become apparent from 12

the following detailed description. It should be understood, however, that the detailed description and the specific examples, while indicating specific embodiments, are given by way of illustration only, since various changes and modifications within the spirit and scope of the present disclosure will become apparent to those skilled in the art from this detailed description. The section headings used herein are for organizational purposes only and are not to be construed as limiting the subject matter described. All documents, or portions of documents, cited in the application including, but not limited to, patents, patent applications, articles, books, manuals, and treatises are hereby expressly incorporated by reference in their entirety for any purpose.

#### INCORPORATION BY REFERENCE

All publications and patent applications mentioned in this specification are herein incorporated by reference to the extent applicable and relevant.

#### BRIEF DESCRIPTION OF THE FIGURES

FIG. 1 shows mean plasma concentration-time profiles of ibrutinib following single oral dose administration of a capsule formulation versus three different wet tablet formulations to fasted beagle dogs (Dose=140 mg).

FIG. 2 shows mean plasma concentration-time profiles of ibrutinib following single oral dose administration of a capsule formulation versus two different dry tablet formulations to fasted beagle dogs (Dose=140 mg).

FIG. 3 is a photo of examples: (A) a capsule comprising 140 mg ibrutinib (Formulation A), and tablets of the invention (B-E) designed to comprise 560 mg, 420 mg, 280 mg, and 140 mg of ibrutinib, respectively.

#### DETAILED DESCRIPTION OF THE INVENTION

The diverse roles played by Btk signaling in various amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin- 40 hematopoietic cell functions, e.g., B-cell receptor activation, suggests that small molecule Btk inhibitors, such as Compound 1, are useful for reducing the risk of or treating a variety of diseases affected by or affecting many cell types of the hematopoietic lineage including, e.g., autoimmune diseases, heteroimmune conditions or diseases, inflammatory diseases, cancer (e.g., B-cell proliferative disorders), and thromboembolic disorders. Further, irreversible Btk inhibitor compounds, such as Compound 1, can be used to inhibit a small subset of other tyrosine kinases that share homology with Btk by having a cysteine residue (including a Cys 481 residue) that can form a covalent bond with the irreversible inhibitor.

In some embodiments, the compositions or tablet formulations comprising Compound 1 can be used in the treatment of an autoimmune disease in a mammal, which includes, but is not limited to, rheumatoid arthritis, psoriatic arthritis, osteoarthritis, Still's disease, juvenile arthritis, lupus, diabetes, myasthenia gravis, Hashimoto's thyroiditis, Ord's thyroiditis, Graves' disease Sjögren's syndrome, multiple sclerosis, Guillain-Barrd syndrome, acute disseminated encephalomyelitis, Addison's disease, opsoclonus-myoclonus syndrome, ankylosing spondylitisis, antiphospholipid antibody syndrome, aplastic anemia, autoimmune hepatitis, coeliac disease, Goodpasture's syndrome, idiopathic thrombocytopenic purpura, optic neuritis, scleroderma, primary biliary cirrhosis, Reiter's syndrome, Takayasu's arteritis, temporal arteritis, warm autoimmune hemolytic anemia,

- - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - - - , - -

Wegener's granulomatosis, psoriasis, alopecia universalis, Behçet's disease, chronic fatigue, dysautonomia, endometriosis, interstitial cystitis, neuromyotonia, scleroderma, and vulvodynia.

13

In some embodiments, the compositions or tablet formulations comprising Compound 1 can be used in the treatment of a heteroimmune disease or condition in a mammal, which include, but are not limited to graft versus host disease, transplantation, transfusion, anaphylaxis, allergies (e.g., allergies to plant pollens, latex, drugs, foods, insect poisons, 10 animal hair, animal dander, dust mites, or cockroach calyx), type I hypersensitivity, allergic conjunctivitis, allergic rhinitis, and atopic dermatitis.

In some embodiments, the compositions or tablet formulations comprising Compound 1 can be used in the treatment 15 of an inflammatory disease in a mammal, which includes, but is not limited to asthma, inflammatory bowel disease, appendicitis, blepharitis, bronchiolitis, bronchitis, bursitis, cervicitis, cholangitis, cholecystitis, colitis, conjunctivitis, dacryoadenitis, dermatitis, dermatomyositis, 20 encephalitis, endocarditis, endometritis, enteritis, enterocolitis, epicondylitis, epididymitis, fasciitis, fibrositis, gastritis, gastroenteritis, hepatitis, hidradenitis suppurativa, laryngitis, mastitis, meningitis, myelitis myocarditis, myositis, nephritis, oophoritis, orchitis, osteitis, otitis, pancreatitis, 25 parotitis, pericarditis, peritonitis, pharyngitis, pleuritis, phlebitis, pneumonitis, pneumonia, proctitis, prostatitis, pyelonephritis, rhinitis, salpingitis, sinusitis, stomatitis, synovitis, tendonitis, tonsillitis, uveitis, vaginitis, vasculitis, and vulvitis. In some embodiments, the inflammatory disease is 30 asthma, appendicitis, blepharitis, bronchiolitis, bronchitis, bursitis, cervicitis, cholangitis, cholecystitis, colitis, conjunctivitis, cystitis, dacryoadenitis, dermatitis, dermatomyositis, encephalitis, endocarditis, endometritis, enteritis, enterocolitis, epicondylitis, epididymitis, fasciitis, fibrositis, 35 gastritis, gastroenteritis, hepatitis, hidradenitis suppurativa, laryngitis, mastitis, meningitis, myelitis myocarditis, myositis, nephritis, oophoritis, orchitis, osteitis, otitis, pancreatitis, parotitis, pericarditis, peritonitis, pharyngitis, pleuritis, phlebitis, pneumonitis, pneumonia, proctitis, prostatitis, 40 pyelonephritis, rhinitis, salpingitis, sinusitis, stomatitis, synovitis, tendonitis, tonsillitis, uveitis, vaginitis, vasculitis, or vulvitis. In some embodiments, the autoimmune disease is inflammatory bowel disease, arthritis, lupus, rheumatoid arthritis, psoriatic arthritis, osteoarthritis, Still's disease, 45 juvenile arthritis, diabetes, myasthenia gravis, Hashimoto's thyroiditis, Ord's thyroiditis, Graves' disease Sjögren's syndrome, multiple sclerosis, Guillain-Barre syndrome, acute disseminated encephalomyelitis, Addison's disease, opsoclonus-myoclonus syndrome, ankylosing spondylitisis, 50 antiphospholipid antibody syndrome, aplastic anemia, autoimmune hepatitis, coeliac disease, Goodpasture's syndrome, idiopathic thrombocytopenic purpura, optic neuritis, scleroderma, primary biliary cirrhosis, Reiter's syndrome, Takayasu's arteritis, temporal arteritis, warm autoimmune hemo- 55 lytic anemia, Wegener's granulomatosis, psoriasis, alopecia universalis, Behçet's disease, chronic fatigue, dysautonomia, endometriosis, interstitial cystitis, neuromyotonia, scleroderma, or vulvodynia.

In yet other embodiments, the methods described herein 60 can be used to treat a cancer, e.g., B-cell proliferative disorders, which include, but are not limited to diffuse large B cell lymphoma, follicular lymphoma, chronic lymphocytic lymphoma, chronic lymphocytic leukemia, B-cell prolymphocytic leukemia, lymphoplasmacytic lymphoma/ 65 Waldenstrom macroglobulinemia, splenic marginal zone lymphoma, plasma cell myeloma, plasmacytoma, extrano-

14

dal marginal zone B cell lymphoma, nodal marginal zone B cell lymphoma, mantle cell lymphoma, mediastinal (thymic) large B cell lymphoma, intravascular large B cell lymphoma, primary effusion lymphoma, burkitt lymphoma/leukemia, and lymphomatoid granulomatosis.

In further embodiments, the methods described herein can be used to treat thromboembolic disorders, which include, but are not limited to myocardial infarct, angina pectoris (including unstable angina), reocclusions or restenoses after angioplasty or aortocoronary bypass, stroke, transitory ischemia, peripheral arterial occlusive disorders, pulmonary embolisms, and deep venous thromboses.

Hematological Malignancies Disclosed herein, in certain em

Disclosed herein, in certain embodiments, is a method for treating a hematological malignancy in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1.

In some embodiments, the hematological malignancy is a non-Hodgkin's lymphoma (NHL). In some embodiments, the hematological malignancy is a chronic lymphocytic leukemia (CLL), small lymphocytic lymphoma (SLL), high risk CLL, or a non-CLL/SLL lymphoma. In some embodiments, the hematological malignancy is follicular lymphoma (FL), diffuse large B-cell lymphoma (DLBCL), mantle cell lymphoma (MCL), Waldenstrom's macroglobulinemia, multiple myeloma (MM), marginal zone lymphoma, Burkitt's lymphoma, non-Burkitt high grade B cell lymphoma, or extranodal marginal zone B cell lymphoma. In some embodiments, the hematological malignancy is acute or chronic myelogenous (or myeloid) leukemia, myelodysplastic syndrome, acute lymphoblastic leukemia, or precursor B-cell acute lymphoblastic leukemia. In some embodiments, the hematological malignancy is chronic lymphocytic leukemia (CLL). In some embodiments, the hematological malignancy is mantle cell lymphoma (MCL). In some embodiments, the hematological malignancy is diffuse large B-cell lymphoma (DLBCL). In some embodiments, the hematological malignancy is diffuse large B-cell lymphoma (DLBCL), ABC subtype. In some embodiments, the hematological malignancy is diffuse large B-cell lymphoma (DLBCL), GCB subtype. In some embodiments, the hematological malignancy is Waldenstrom's macroglobulinemia (WM). In some embodiments, the hematological malignancy is multiple myeloma (MM). In some embodiments, the hematological malignancy is Burkitt's lymphoma. In some embodiments, the hematological malignancy is follicular lymphoma (FL). In some embodiments, the hematological malignancy is transformed follicular lymphoma. In some embodiments, the hematological malignancy is marginal zone lymphoma.

In some embodiments, the hematological malignancy is relapsed or refractory non-Hodgkin's lymphoma (NHL). In some embodiments, the hematological malignancy is relapsed or refractory diffuse large B-cell lymphoma (DLBCL), relapsed or refractory mantle cell lymphoma (MCL), relapsed or refractory follicular lymphoma (FL), relapsed or refractory CLL, relapsed or refractory SLL, relapsed or refractory multiple myeloma, relapsed or refractory Waldenstrom's macroglobulinemia, relapsed or refractory multiple myeloma (MM), relapsed or refractory marginal zone lymphoma, relapsed or refractory Burkitt's lymphoma, relapsed or refractory non-Burkitt high grade B cell lymphoma, relapsed or refractory extranodal marginal zone B cell lymphoma. In some embodiments, the hematological malignancy is a relapsed or refractory acute or chronic myelogenous (or myeloid) leukemia, relapsed or

refractory myelodysplastic syndrome, relapsed or refractory acute lymphoblastic leukemia, or relapsed or refractory precursor B-cell acute lymphoblastic leukemia. In some embodiments, the hematological malignancy is relapsed or refractory chronic lymphocytic leukemia (CLL). In some 5 embodiments, the hematological malignancy is relapsed or refractory mantle cell lymphoma (MCL). In some embodiments, the hematological malignancy is relapsed or refractory diffuse large B-cell lymphoma (DLBCL). In some embodiments, the hematological malignancy is relapsed or refractory diffuse large B-cell lymphoma (DLBCL), ABC subtype. In some embodiments, the hematological malignancy is relapsed or refractory diffuse large B-cell lymphoma (DLBCL), GCB subtype. In some embodiments, the hematological malignancy is relapsed or refractory Walden- 15 strom's macroglobulinemia (WM). In some embodiments, the hematological malignancy is relapsed or refractory multiple myeloma (MM). In some embodiments, the hematological malignancy is relapsed or refractory Burkitt's lymphoma. In some embodiments, the hematological 20 malignancy is relapsed or refractory follicular lymphoma (FL).

In some embodiments, the hematological malignancy is a hematological malignancy that is classified as high-risk. In some embodiments, the hematological malignancy is high 25 risk CLL or high risk SLL.

B-cell lymphoproliferative disorders (BCLDs) are neoplasms of the blood and encompass, inter alia, non-Hodgkin lymphoma, multiple myeloma, and leukemia. BCLDs can originate either in the lymphatic tissues (as in the case of 30 lymphoma) or in the bone marrow (as in the case of leukemia and myeloma), and they all are involved with the uncontrolled growth of lymphocytes or white blood cells. There are many subtypes of BCLD, e.g., chronic lymphocytic leukemia (CLL) and non-Hodgkin lymphoma (NHL). 35 The disease course and treatment of BCLD is dependent on the BCLD subtype; however, even within each subtype the clinical presentation, morphologic appearance, and response to therapy is heterogeneous.

Malignant lymphomas are neoplastic transformations of 40 cells that reside predominantly within lymphoid tissues. Two groups of malignant lymphomas are Hodgkin's lymphoma and non-Hodgkin's lymphoma (NHL). Both types of lymphomas infiltrate reticuloendothelial tissues. However, they differ in the neoplastic cell of origin, site of disease, presence 45 of systemic symptoms, and response to treatment (Freedman et al., "Non-Hodgkin's Lymphomas" Chapter 134, Cancer Medicine, (an approved publication of the American Cancer Society, B.C. Decker Inc., Hamilton, Ontario, 2003).

Non-Hodgkin's Lymphomas

Disclosed herein, in certain embodiments, is a method for treating a non-Hodgkin's lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1.

Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory non-Hodgkin's lymphoma in an individual in need thereof, comprising: administering to the individual a therapeutically-effective amount of Compound 1. In some embodiments, the non-Hodgkin's lymphoma is relapsed or refractory diffuse large B-cell lymphoma (DLBCL), relapsed or refractory mantle cell lymphoma, relapsed or refractory follicular lymphoma, or relapsed or refractory CLL.

Non-Hodgkin lymphomas (NHL) are a diverse group of 65 malignancies that are predominately of B-cell origin. NHL may develop in any organs associated with lymphatic sys-

tem such as spleen, lymph nodes or tonsils and can occur at any age. NHL is often marked by enlarged lymph nodes, fever, and weight loss. NHL is classified as either B-cell or T-cell NHL. Lymphomas related to lymphoproliferative disorders following bone marrow or stem cell transplantation are usually B-cell NHL. In the Working Formulation classification scheme, NHL has been divided into low-, intermediate-, and high-grade categories by virtue of their natural histories (see "The Non-Hodgkin's Lymphoma Pathologic Classification Project," Cancer 49(1982):2112-2135). The low-grade lymphomas are indolent, with a median survival of 5 to 10 years (Homing and Rosenberg (1984) N. Engl. J. Med. 311:1471-1475). Although chemotherapy can induce remissions in the majority of indolent lymphomas, cures are rare and most patients eventually relapse, requiring further therapy. The intermediate- and high-grade lymphomas are more aggressive tumors, but they have a greater chance for cure with chemotherapy. However, a significant proportion of these patients will relapse and require further treatment.

16

A non-limiting list of the B-cell NHL includes Burkitt's lymphoma (e.g., Endemic Burkitt's Lymphoma and Sporadic Burkitt's Lymphoma), Cutaneous B-Cell Lymphoma, Cutaneous Marginal Zone Lymphoma (MZL), Diffuse Large Cell Lymphoma (DLBCL), Diffuse Mixed Small and Large Cell Lymphoma, Diffuse Small Cleaved Cell, Diffuse Small Lymphocytic Lymphoma, Extranodal Marginal Zone B-cell lymphoma, follicular lymphoma, Follicular Small Cleaved Cell (Grade 1), Follicular Mixed Small Cleaved and Large Cell (Grade 2), Follicular Large Cell (Grade 3), Intravascular Large B-Cell Lymphoma, Intravascular Lymphomatosis, Large Cell Immunoblastic Lymphoma, Large Cell Lymphoma (LCL), Lymphoblastic Lymphoma, MALT Lymphoma, Mantle Cell Lymphoma (MCL), immunoblastic large cell lymphoma, precursor B-lymphoblastic lymphoma, mantle cell lymphoma, chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL), extranodal marginal zone B-cell lymphoma-mucosa-associated lymphoid tissue (MALT) lymphoma, Mediastinal Large B-Cell Lymphoma, nodal marginal zone B-cell lymphoma, splenic marginal zone B-cell lymphoma, primary mediastinal B-cell lymphoma, lymphoplasmocytic lymphoma, hairy cell leukemia, Waldenstrom's Macroglobulinemia, and primary central nervous system (CNS) lymphoma. Additional non-Hodgkin's lymphomas are contemplated within the scope of the present invention and apparent to those of ordinary skill in the art.

DLBCL

Disclosed herein, in certain embodiments, is a method for treating a DLCBL in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory DLCBL in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

As used herein, the term "Diffuse large B-cell lymphoma (DLBCL)" refers to a neoplasm of the germinal center B lymphocytes with a diffuse growth pattern and a high-intermediate proliferation index. DLBCLs represent approximately 30% of all lymphomas and may present with several morphological variants including the centroblastic, immunoblastic, T-cell/histiocyte rich, anaplastic and plasmoblastic subtypes. Genetic tests have shown that there are different subtypes of DLBCL. These subtypes seem to have

different outlooks (prognoses) and responses to treatment. DLBCL can affect any age group but occurs mostly in older people (the average age is mid-60s).

Disclosed herein, in certain embodiments, is a method for treating diffuse large B-cell lymphoma, activated B cell-like 5 subtype (ABC-DLBCL), in an individual in need thereof, comprising: administering to the individual an irreversible Btk inhibitor in an amount from 300 mg/day up to, and including, 1000 mg/day. The ABC subtype of diffuse large B-cell lymphoma (ABC-DLBCL) is thought to arise from 10 post germinal center B cells that are arrested during plasmatic differentiation. The ABC subtype of DLBCL (ABC-DLBCL) accounts for approximately 30% total DLBCL diagnoses. It is considered the least curable of the DLBCL molecular subtypes and, as such, patients diagnosed with the 15 ABC-DLBCL typically display significantly reduced survival rates compared with individuals with other types of DLCBL. ABC-DLBCL is most commonly associated with chromosomal translocations deregulating the germinal center master regulator BCL6 and with mutations inactivating 20 the PRDM1 gene, which encodes a transcriptional repressor required for plasma cell differentiation.

A particularly relevant signaling pathway in the pathogenesis of ABC-DLBCL is the one mediated by the nuclear factor (NF)-κB transcription complex. The NF-κB family 25 comprises 5 members (p50, p52, p65, c-rel and RelB) that form homo- and heterodimers and function as transcriptional factors to mediate a variety of proliferation, apoptosis, inflammatory and immune responses and are critical for normal B-cell development and survival. NF-κB is widely used by eukaryotic cells as a regulator of genes that control cell proliferation and cell survival. As such, many different types of human tumors have misregulated NF-κB: that is, NF-κd is constitutively active. Active NF-κd turns on the expression of genes that keep the cell proliferating and 35 protect the cell from conditions that would otherwise cause it to die via apoptosis.

The dependence of ABC DLBCLs on NF-kB depends on a signaling pathway upstream of IkB kinase comprised of CARD11, BCL10 and MALT1 (the CBM complex). Inter- 40 ference with the CBM pathway extinguishes NF-kB signaling in ABC DLBCL cells and induces apoptosis. The molecular basis for constitutive activity of the NF-kB pathway is a subject of current investigation but some somatic alterations to the genome of ABC DLBCLs clearly invoke 45 this pathway. For example, somatic mutations of the coiledcoil domain of CARD11 in DLBCL render this signaling scaffold protein able to spontaneously nucleate proteinprotein interaction with MALT1 and BCL10, causing IKK activity and NF-kB activation. Constitutive activity of the B 50 cell receptor signaling pathway has been implicated in the activation of NF-kB in ABC DLBCLs with wild type CARD11, and this is associated with mutations within the cytoplasmic tails of the B cell receptor subunits CD79A and CD79B. Oncogenic activating mutations in the signaling 55 adapter MYD88 activate NF-kB and synergize with B cell receptor signaling in sustaining the survival of ABC DLBCL cells. In addition, inactivating mutations in a negative regulator of the NF-kB pathway, A20, occur almost exclusively in ABC DLBCL.

Indeed, genetic alterations affecting multiple components of the NF- $\kappa$ B signaling pathway have been recently identified in more than 50% of ABC-DLBCL patients, where these lesions promote constitutive NF- $\kappa$ B activation, thereby contributing to lymphoma growth. These include mutations of 65 CARD11 (~10% of the cases), a lymphocyte-specific cytoplasmic scaffolding protein that—together with MALT1 and

18

BCL10—forms the BCR signalosome, which relays signals from antigen receptors to the downstream mediators of NF-κB activation. An even larger fraction of cases (~30%) carry biallelic genetic lesions inactivating the negative NF-κB regulator A20. Further, high levels of expression of NF-κB target genes have been observed in ABC-DLBCL tumor samples. See, e.g., U. Klein et al., (2008), *Nature Reviews Immunology* 8:22-23; R. E. Davis et al., (2001), Journal of Experimental Medicine 194:1861-1874; G. Lentz et al., (2008), *Science* 319:1676-1679; M. Compagno et al., (2009), *Nature* 459:712-721; and L. Srinivasan et al., (2009), *Cell* 139:573-586).

DLBCL cells of the ABC subtype, such as OCI-Ly10, have chronic active BCR signaling and are very sensitive to the Btk inhibitor described herein. The irreversible Btk inhibitor described herein potently and irreversibly inhibits the growth of OCI-Ly10 (EC<sub>50</sub> continuous exposure=10 nM, EC<sub>50</sub> 1 hour pulse=50 nM). In addition, induction of apoptosis, as shown by capsase activation, Annexin-V flow cytometry and increase in sub-GO fraction is observed in OCILy 10. Both sensitive and resistant cells express Btk at similar levels, and the active site of Btk is fully occupied by the inhibitor in both as shown using a fluorescently labeled affinity probe. OCI-Ly10 cells are shown to have chronically active BCR signaling to NF-kB which is dose dependently inhibited by the Btk inhibitors described herein. The activity of Btk inhibitors in the cell lines studied herein are also characterized by comparing signal transduction profiles (Btk, PLCy, ERK, NF-kB, AKT), cytokine secretion profiles and mRNA expression profiles, both with and without BCR stimulation, and observed significant differences in these profiles that lead to clinical biomarkers that identify the most sensitive patient populations to Btk inhibitor treatment. See U.S. Pat. No. 7,711,492 and Staudt et al., Nature, Vol. 463, Jan. 7, 2010, pp. 88-92, the contents of which are incorporated by reference in their entirety.

Follicular Lymphoma

Disclosed herein, in certain embodiments, is a method for treating a follicular lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory follicular lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

As used herein, the term "follicular lymphoma" refers to any of several types of non-Hodgkin's lymphoma in which the lymphomatous cells are clustered into nodules or follicles. The term follicular is used because the cells tend to grow in a circular, or nodular, pattern in lymph nodes. The average age for people with this lymphoma is about 60.

CLL/SLL

Disclosed herein, in certain embodiments, is a method for treating a CLL or SLL in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory CLL or SLL in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

Chronic lymphocytic leukemia and small lymphocytic lymphoma (CLL/SLL) are commonly thought as the same disease with slightly different manifestations. Where the

cancerous cells gather determines whether it is called CLL or SLL. When the cancer cells are primarily found in the lymph nodes, lima bean shaped structures of the lymphatic system (a system primarily of tiny vessels found in the body), it is called SLL. SLL accounts for about 5% to 10% of all lymphomas. When most of the cancer cells are in the bloodstream and the bone marrow, it is called CLL.

Both CLL and SLL are slow-growing diseases, although CLL, which is much more common, tends to grow slower. CLL and SLL are treated the same way. They are usually not 10 considered curable with standard treatments, but depending on the stage and growth rate of the disease, most patients live longer than 10 years. Occasionally over time, these slow-growing lymphomas may transform into a more aggressive type of lymphoma.

Chronic lymphoid leukemia (CLL) is the most common type of leukemia. It is estimated that 100,760 people in the United States are living with or are in remission from CLL. Most (>75%) people newly diagnosed with CLL are over the age of 50. Currently CLL treatment focuses on controlling 20 the disease and its symptoms rather than on an outright cure. CLL is treated by chemotherapy, radiation therapy, biological therapy, or bone marrow transplantation. Symptoms are sometimes treated surgically (splenectomy removal of enlarged spleen) or by radiation therapy ("de-bulking" swol- 25 len lymph nodes). Though CLL progresses slowly in most cases, it is considered generally incurable. Certain CLLs are classified as high-risk. As used herein, "high risk CLL" means CLL characterized by at least one of the following 1) 17p13-; 2) 11q22-; 3) unmutated IgVH together with ZAP- 30 70+ and/or CD38+; or 4) trisomy 12.

CLL treatment is typically administered when the patient's clinical symptoms or blood counts indicate that the disease has progressed to a point where it may affect the patient's quality of life.

Small lymphocytic leukemia (SLL) is very similar to CLL described supra, and is also a cancer of B-cells. In SLL the abnormal lymphocytes mainly affect the lymph nodes. However, in CLL the abnormal cells mainly affect the blood and the bone marrow. The spleen may be affected in both 40 conditions. SLL accounts for about 1 in 25 of all cases of non-Hodgkin lymphoma. It can occur at any time from young adulthood to old age, but is rare under the age of 50. SLL is considered an indolent lymphoma. This means that the disease progresses very slowly, and patients tend to live 45 many years after diagnosis. However, most patients are diagnosed with advanced disease, and although SLL responds well to a variety of chemotherapy drugs, it is generally considered to be incurable. Although some cancers tend to occur more often in one gender or the other, cases 50 and deaths due to SLL are evenly split between men and women. The average age at the time of diagnosis is 60 years.

Although SLL is indolent, it is persistently progressive. The usual pattern of this disease is one of high response rates to radiation therapy and/or chemotherapy, with a period of 55 disease remission. This is followed months or years later by an inevitable relapse. Re-treatment leads to a response again, but again the disease will relapse. This means that although the short-term prognosis of SLL is quite good, over time, many patients develop fatal complications of recurrent disease. Considering the age of the individuals typically diagnosed with CLL and SLL, there is a need in the art for a simple and effective treatment of the disease with minimum side-effects that do not impede on the patient's quality of life. The instant invention fulfills this long standing need in 65 the art.

Mantle Cell Lymphoma

20

Disclosed herein, in certain embodiments, is a method for treating a Mantle cell lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory Mantle cell lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

As used herein, the term, "Mantle cell lymphoma" refers to a subtype of B-cell lymphoma, due to CD5 positive antigen-naive pregerminal center B-cell within the mantle zone that surrounds normal germinal center follicles. MCL cells generally over-express cyclin D1 due to a t(11:14) chromosomal translocation in the DNA. More specifically, the translocation is at t(11;14)(q13;q32). Only about 5% of lymphomas are of this type. The cells are small to medium in size. Men are affected most often. The average age of patients is in the early 60s. The lymphoma is usually widespread when it is diagnosed, involving lymph nodes, bone marrow, and, very often, the spleen. Mantle cell lymphoma is not a very fast growing lymphoma, but is difficult to treat.

Marginal Zone B-Cell Lymphoma

Disclosed herein, in certain embodiments, is a method for treating a marginal zone B-cell lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory marginal zone B-cell lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

As used herein, the term "marginal zone B-cell lymphoma" refers to a group of related B-cell neoplasms that involve the lymphoid tissues in the marginal zone, the patchy area outside the follicular mantle zone. Marginal zone lymphomas account for about 5% to 10% of lymphomas. The cells in these lymphomas look small under the microscope. There are 3 main types of marginal zone lymphomas, nodal marginal zone B-cell lymphomas, nodal marginal zone B-cell lymphoma, and splenic marginal zone lymphoma.

MALT

Disclosed herein, in certain embodiments, is a method for treating a MALT in an individual in need thereof, comprising: administering to the individual an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory MALT in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

The term "mucosa-associated lymphoid tissue (MALT) lymphoma", as used herein, refers to extranodal manifestations of marginal-zone lymphomas. Most MALT lymphoma are a low grade, although a minority either manifest initially as intermediate-grade non-Hodgkin lymphoma (NHL) or evolve from the low-grade form. Most of the MALT lymphoma occur in the stomach, and roughly 70% of gastric MALT lymphoma are associated with *Helicobacter pylori* infection. Several cytogenetic abnormalities have been identified, the most common being trisomy 3 or t(11;18). Many of these other MALT lymphoma have also been linked to

infections with bacteria or viruses. The average age of patients with MALT lymphoma is about 60.

Nodal Marginal Zone B-Cell Lymphoma

Disclosed herein, in certain embodiments, is a method for treating a nodal marginal zone B-cell lymphoma in an 5 individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory nodal marginal zone B-cell lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

The term "nodal marginal zone B-cell lymphoma" refers 15 to an indolent B-cell lymphoma that is found mostly in the lymph nodes. The disease is rare and only accounts for 1% of all Non-Hodgkin's Lymphomas (NHL). It is most commonly diagnosed in older patients, with women more susceptible than men. The disease is classified as a marginal 20 zone lymphoma because the mutation occurs in the marginal zone of the B-cells. Due to its confinement in the lymph nodes, this disease is also classified as nodal.

Splenic Marginal Zone B-Cell Lymphoma

Disclosed herein, in certain embodiments, is a method for 25 treating a splenic marginal zone B-cell lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for 30 treating relapsed or refractory splenic marginal zone B-cell lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

The term "splenic marginal zone B-cell lymphoma" refers to specific low-grade small B-cell lymphoma that is incorporated in the World Health Organization classification. Characteristic features are splenomegaly, moderate lymphocytosis with villous morphology, intrasinusoidal pattern of 40 involvement of various organs, especially bone marrow, and relative indolent course. Tumor progression with increase of blastic forms and aggressive behavior are observed in a minority of patients. Molecular and cytogenetic studies have shown heterogeneous results probably because of the lack of 45 standardized diagnostic criteria.

Burkitt Lymphoma

Disclosed herein, in certain embodiments, is a method for treating a Burkitt lymphoma in an individual in need thereof, comprising: administering to the individual a composition or 50 tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory Burkitt lymphoma in an individual in need thereof, comprising: administering to the individual a composition or tablet 55 formulation described herein comprising a therapeutically-effective amount of Compound 1.

The term "Burkitt lymphoma" refers to a type of Non-Hodgkin Lymphoma (NHL) that commonly affects children. It is a highly aggressive type of B-cell lymphoma that often 60 starts and involves body parts other than lymph nodes. In spite of its fast-growing nature, Burkitt's lymphoma is often curable with modern intensive therapies. There are two broad types of Burkitt's lymphoma—the sporadic and the endemic varieties:

Endemic Burkitt's lymphoma: The disease involves children much more than adults, and is related to Epstein Barr Virus (EBV) infection in 95% cases. It occurs primarily is equatorial Africa, where about half of all childhood cancers are Burkitt's lymphoma. It characteristically has a high chance of involving the jawbone, a rather distinctive feature that is rare in sporadic Burkitt's. It also commonly involves the abdomen.

22

Sporadic Burkitt's lymphoma: The type of Burkitt's lymphoma that affects the rest of the world, including Europe and the Americas is the sporadic type. Here too, it's mainly a disease in children. The link between Epstein Barr Virus (EBV) is not as strong as with the endemic variety, though direct evidence of EBV infection is present in one out of five patients. More than the involvement of lymph nodes, it is the abdomen that is notably affected in more than 90% of the children. Bone marrow involvement is more common than in the sporadic variety.

Waldenstrom Macroglobulinemia

Disclosed herein, in certain embodiments, is a method for treating a Waldenstrom macroglobulinemia in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory Waldenstrom macroglobulinemia in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

The term "Waldenstrom macroglobulinemia", also known as lymphoplasmacytic lymphoma, is cancer involving a subtype of white blood cells called lymphocytes. It is characterized by an uncontrolled clonal proliferation of terminally differentiated B lymphocytes. It is also characterized by the lymphoma cells making an antibody called immunoglobulin M (IgM). The IgM antibodies circulate in the blood in large amounts, and cause the liquid part of the blood to thicken, like syrup. This can lead to decreased blood flow to many organs, which can cause problems with vision (because of poor circulation in blood vessels in the back of the eyes) and neurological problems (such as headache, dizziness, and confusion) caused by poor blood flow within the brain. Other symptoms can include feeling tired and weak, and a tendency to bleed easily. The underlying etiology is not fully understood but a number of risk factors have been identified, including the locus 6p21.3 on chromosome 6. There is a 2- to 3-fold risk increase of developing WM in people with a personal history of autoimmune diseases with autoantibodies and particularly elevated risks associated with hepatitis, human immunodeficiency virus, and rickettsiosis.

Multiple Myeloma

Disclosed herein, in certain embodiments, is a method for treating a myeloma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory myeloma in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective amount of Compound 1.

Multiple myeloma, also known as MM, myeloma, plasma cell myeloma, or as Kahler's disease (after Otto Kahler) is a cancer of the white blood cells known as plasma cells. A type of B cell, plasma cells are a crucial part of the immune system responsible for the production of antibodies in humans and other vertebrates. They are produced in the bone marrow and are transported through the lymphatic system.

Leukemia

Disclosed herein, in certain embodiments, is a method for treating a leukemia in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising an amount of 5 Compound 1. Further disclosed herein, in certain embodiments, is a method for treating relapsed or refractory leukemia in an individual in need thereof, comprising: administering to the individual a composition or tablet formulation described herein comprising a therapeutically-effective 10 amount of Compound 1.

Leukemia is a cancer of the blood or bone marrow characterized by an abnormal increase of blood cells, usually leukocytes (white blood cells). Leukemia is a broad term covering a spectrum of diseases. The first division is 15 between its acute and chronic forms: (i) acute leukemia is characterized by the rapid increase of immature blood cells. This crowding makes the bone marrow unable to produce healthy blood cells. Immediate treatment is required in acute leukemia due to the rapid progression and accumulation of 20 the malignant cells, which then spill over into the bloodstream and spread to other organs of the body. Acute forms of leukemia are the most common forms of leukemia in children; (ii) chronic leukemia is distinguished by the excessive build up of relatively mature, but still abnormal, white 25 blood cells. Typically taking months or years to progress, the cells are produced at a much higher rate than normal cells, resulting in many abnormal white blood cells in the blood. Chronic leukemia mostly occurs in older people, but can theoretically occur in any age group. Additionally, the 30 diseases are subdivided according to which kind of blood cell is affected. This split divides leukemias into lymphoblastic or lymphocytic leukemias and myeloid or myelogenous leukemias: (i) lymphoblastic or lymphocytic leukemias, the cancerous change takes place in a type of marrow 35 cell that normally goes on to form lymphocytes, which are infection-fighting immune system cells; (ii) myeloid or myelogenous leukemias, the cancerous change takes place in a type of marrow cell that normally goes on to form red blood cells, some other types of white cells, and platelets. 40

Within these main categories, there are several subcategories including, but not limited to, Acute lymphoblastic leukemia (ALL), precursor B-cell acute lymphoblastic leukemia (precursor B-ALL; also called precursor B-lymphoblastic leukemia), Acute myelogenous leukemia (AML), 45 Chronic myelogenous leukemia (CML), and Hairy cell leukemia (HCL). Accordingly, disclosed herein, in certain embodiments, is a method for treating Acute lymphoblastic leukemia (ALL), precursor B-cell acute lymphoblastic leukemia (precursor B-ALL; also called precursor B-lympho- 50 blastic leukemia), Acute myelogenous leukemia (AML), Chronic myelogenous leukemia (CML), or Hairy cell leukemia (HCL) in an individual in need thereof, comprising: administering to the individual an amount of Compound 1. In some embodiments, the leukemia is a relapsed or refrac- 55 tory leukemia. In some embodiments, the leukemia is a relapsed or refractory Acute lymphoblastic leukemia (ALL), relapsed or refractory precursor B-cell acute lymphoblastic leukemia (precursor B-ALL; also called precursor B-lymphoblastic leukemia), relapsed or refractory Acute myelog- 60 enous leukemia (AML), relapsed or refractory Chronic myelogenous leukemia (CML), or relapsed or refractory Hairy cell leukemia (HCL).

Symptoms, diagnostic tests, and prognostic tests for each of the above-mentioned conditions are known. See, e.g., 65 *Harrison's Principles of Internal Medicine*©," 16th ed., 2004, The McGraw-Hill Companies, Inc. Dey et al. (2006),

24

Cytojournal 3(24), and the "Revised European American Lymphoma" (REAL) classification system (see, e.g., the website maintained by the National Cancer Institute).

A number of animal models of are useful for establishing a range of therapeutically effective doses of irreversible Btk inhibitor compounds, such as Compound 1, for treating any of the foregoing diseases.

The therapeutic efficacy of Compound 1 for any one of the foregoing diseases can be optimized during a course of treatment. For example, a subject being treated can undergo a diagnostic evaluation to correlate the relief of disease symptoms or pathologies to inhibition of in vivo Btk activity achieved by administering a given dose of Compound 1. Cellular assays known in the art can be used to determine in vivo activity of Btk in the presence or absence of an irreversible Btk inhibitor. For example, since activated Btk is phosphorylated at tyrosine 223 (Y223) and tyrosine 551 (Y551), phospho-specific immunocytochemical staining of P-Y223 or P-Y551-positive cells can be used to detect or quantify activation of Btk in a population of cells (e.g., by FACS analysis of stained vs unstained cells). See, e.g., Nisitani et al. (1999), Proc. Natl. Acad. Sci, USA 96:2221-2226. Thus, the amount of the Btk inhibitor compound that is administered to a subject can be increased or decreased as needed so as to maintain a level of Btk inhibition optimal for treating the subject's disease state.

Compound 1 can irreversibly inhibit Btk and may be used to treat mammals suffering from Bruton's tyrosine kinase-dependent or Bruton's tyrosine kinase mediated conditions or diseases, including, but not limited to, cancer, autoimmune and other inflammatory diseases. Compound 1 has shown efficacy is a wide variety of diseases and conditions that are described herein.

In some embodiments, Compound 1 is used for the manufacture of a medicament for treating any of the foregoing conditions (e.g., autoimmune diseases, inflammatory diseases, allergy disorders, B-cell proliferative disorders, or thromboembolic disorders).

Compound 1, and Pharmaceutically Acceptable Salts Thereof

The Btk inhibitor compound described herein (i.e. Compound 1) is selective for Btk and kinases having a cysteine residue in an amino acid sequence position of the tyrosine kinase that is homologous to the amino acid sequence position of cysteine 481 in Btk. The Btk inhibitor compound can form a covalent bond with Cys 481 of Btk (e.g., via a Michael reaction).

"Compound 1" or "1-((R)-3-(4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl)piperidin-1-yl) prop-2-en-1-one" or "1-{(3R)-3-[4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl]piperidin-1-yl}prop-2-en-1-one" or "2-Propen-1-one, 1-[(3R)-3-[4-amino-3-(4-phenoxyphenyl)-1H-pyrazolo[3,4-d]pyrimidin-1-yl]-1-piperidinyl-" or ibrutinib or any other suitable name refers to the compound with the following structure:

A wide variety of pharmaceutically acceptable salts is formed from Compound 1 and includes:

acid addition salts formed by reacting Compound 1 with an organic acid, which includes aliphatic mono- and dicarboxylic acids, phenyl-substituted alkanoic acids, 25 hydroxyl alkanoic acids, alkanedioic acids, aromatic acids, aliphatic and aromatic sulfonic acids, amino acids, etc. and include, for example, acetic acid, trifluoroacetic acid, propionic acid, glycolic acid, pyruvic acid, oxalic acid, maleic acid, malonic acid, succinic 30 acid, fumaric acid, tartaric acid, citric acid, benzoic acid, cinnamic acid, mandelic acid, methanesulfonic acid, ethanesulfonic acid, p-toluenesulfonic acid, salicylic acid, and the like;

acid addition salts formed by reacting Compound 1 with 35 an inorganic acid, which includes hydrochloric acid, hydrobromic acid, sulfuric acid, nitric acid, phosphoric acid, hydroiodic acid, hydrofluoric acid, phosphorous acid, and the like.

The term "pharmaceutically acceptable salts" in reference 40 to Compound 1 refers to a salt of Compound 1, which does not cause significant irritation to a mammal to which it is administered and does not substantially abrogate the biological activity and properties of the compound.

It should be understood that a reference to a pharmaceu- 45 tically acceptable salt includes the solvent addition forms (solvates). Solvates contain either stoichiometric or nonstoichiometric amounts of a solvent, and are formed during the process of product formation or isolation with pharmaceutically acceptable solvents such as water, ethanol, metha- 50 nol, methyl tert-butyl ether (MTBE), diisopropyl ether (DIPE), ethyl acetate, isopropyl acetate, isopropyl alcohol, methyl isobutyl ketone (MIBK), methyl ethyl ketone (MEK), acetone, nitromethane, tetrahydrofuran (THF), dichloromethane (DCM), dioxane, heptanes, toluene, ani- 55 sole, acetonitrile, and the like. In one aspect, solvates are formed using, but not limited to, Class 3 solvent(s). Categories of solvents are defined in, for example, the International Conference on Harmonization of Technical Requirements for Registration of Pharmaceuticals for Human Use 60 (ICH), "Impurities: Guidelines for Residual Solvents, Q3C (R3), (November 2005). Hydrates are formed when the solvent is water, or alcoholates are formed when the solvent is alcohol. In some embodiments, solvates of Compound 1, or pharmaceutically acceptable salts thereof, are conve- 65 niently prepared or formed during the processes described herein. In some embodiments, solvates of Compound 1 are

anhydrous. In some embodiments, Compound 1, or pharmaceutically acceptable salts thereof, exist in unsolvated form. In some embodiments, Compound 1, or pharmaceutically acceptable salts thereof, exist in unsolvated form and are anhydrous.

In yet other embodiments, Compound 1, or a pharmaceutically acceptable salt thereof, is prepared in various forms, including but not limited to, amorphous phase, crystalline forms, milled forms and nano-particulate forms. In some embodiments, Compound 1, or a pharmaceutically acceptable salt thereof, is amorphous and embodiments, Compound 1, or a pharmaceutically acceptable salt thereof, is amorphous and anhydrous. In some embodiments, Compound 1, or a pharmaceutically acceptable salt thereof, is crystalline. In some embodiments, Compound 1, or a pharmaceutically acceptable salt thereof, is crystalline and anhydrous.

In some embodiments, Compound 1 is prepared as out-  $_{20}$  lined in U.S. Pat. No. 7,514,444.

#### Certain Terminology

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as is commonly understood by one of skill in the art to which the claimed subject matter belongs. It is to be understood that the foregoing general description and the following detailed description are exemplary and explanatory only and are not restrictive of any subject matter claimed. In this application, the use of the singular includes the plural unless specifically stated otherwise. It must be noted that, as used in the specification and the appended claims, the singular forms "a," "an" and "the" include plural referents unless the context clearly dictates otherwise. In this application, the use of "or" means "and/or" unless stated otherwise. Furthermore, use of the term "including" as well as other forms, such as "include", "includes," and "included," is not limiting.

The section headings used herein are for organizational purposes only and are not to be construed as limiting the subject matter described. All documents, or portions of documents, cited in the application including, but not limited to, patents, patent applications, articles, books, manuals, and treatises are hereby expressly incorporated by reference in their entirety for any purpose.

The term "about" when used before a numerical value indicates that the value may vary within a reasonable range, such as within +10%. +5% or +1% of the stated value.

As used herein, the term "comprising" or its grammatic variants is intended to mean that the compositions and methods, etc., include the recited elements, but do not exclude others. "Consisting essentially of" or its grammatic variants when used to define compositions and methods, shall mean excluding other elements of any essential significance to the combination for the intended use, but not excluding elements that do not materially affect the characteristic(s) of the compositions or methods. "Consisting of" or its grammatic variants shall mean excluding elements not specifically recited. Embodiments defined by each of these transition terms are within the scope of this invention. For example, when a formulation is described as comprising ingredients A, B and C, a formulation consisting essentially of A, B and C, and a formulation consisting of A, B and C are independently within the scope of this invention.

The term "acceptable" or "pharmaceutically acceptable", with respect to a formulation, composition or ingredient, as used herein, means having no persistent detrimental effect on the general health of the subject being treated or does not

abrogate the biological activity or properties of the compound, and is relatively nontoxic.

As used herein, the term "agonist" refers to a compound, the presence of which results in a biological activity of a protein that is the same as the biological activity resulting 5 from the presence of a naturally occurring ligand for the protein, such as, for example, Btk.

As used herein, the term "partial agonist" refers to a compound the presence of which results in a biological activity of a protein that is of the same type as that resulting from the presence of a naturally occurring ligand for the protein, but of a lower magnitude.

As used herein, the term "antagonist" refers to a compound, the presence of which results in a decrease in the magnitude of a biological activity of a protein. In certain 15 embodiments, the presence of an antagonist results in complete inhibition of a biological activity of a protein, such as, for example, Btk. In certain embodiments, an antagonist is an inhibitor.

As used herein, "amelioration" of the symptoms of a 20 particular disease, disorder or condition by administration of a particular compound or pharmaceutical composition refers to any lessening of severity, delay in onset, slowing of progression, or shortening of duration, whether permanent or temporary, lasting or transient that can be attributed to or 25 associated with administration of the compound or composition.

"Bioavailability" refers to the percentage of Compound 1 dosed that is delivered into the general circulation of the animal or human being studied. The total exposure (AUC  $_{(0-\infty)}$ ) of a drug when administered intravenously is usually defined as 100% bioavailable (F %). "Oral bioavailability" refers to the extent to which Compound 1 is absorbed into the general circulation when the pharmaceutical composition is taken orally as compared to intravenous injection.

"Blood plasma concentration" refers to the concentration of Compound 1 in the plasma component of blood of a subject. It is understood that the plasma concentration of Compound 1 may vary significantly between subjects, due to variability with respect to metabolism and/or possible 40 interactions with other therapeutic agents. In accordance with one embodiment disclosed herein, the blood plasma concentration of Compound 1 may vary from subject to subject. Likewise, values such as maximum plasma concentration ( $C_{max}$ ) or time to reach maximum plasma concentration ( $T_{max}$ ), or total area under the plasma concentration time curve ( $AUC_{(O-\infty)}$ ) may vary from subject to subject. Due to this variability, the amount necessary to constitute "a therapeutically effective amount" of Compound 1 may vary from subject to subject.

The term "Bruton's tyrosine kinase," as used herein, refers to Bruton's tyrosine kinase from *Homo sapiens*, as disclosed in, e.g., U.S. Pat. No. 6,326,469 (GenBank Accession No. NP\_000052).

The terms "co-administration" or the like, as used herein, 55 are meant to encompass administration of the selected therapeutic agents to a single patient, and are intended to include treatment regimens in which the agents are administered by the same or different route of administration or at the same or different time. In some embodiments, the term "co-administration" or the like, is meant to encompass the administration of the selected therapeutic agents in the same cycle(s). In these embodiments, the selected therapeutic agents may be administered on the same or different days of the cycle(s).

The terms "effective amount" or "therapeutically effective amount," as used herein, refer to a sufficient amount of an

28

agent or a compound being administered which will relieve to some extent one or more of the symptoms of the disease or condition being treated. The result can be reduction and/or alleviation of the signs, symptoms, or causes of a disease, or any other desired alteration of a biological system. For example, an "effective amount" for therapeutic uses is the amount of the composition including a compound as disclosed herein required to provide a clinically significant decrease in disease symptoms without undue adverse side effects. An appropriate "effective amount" in any individual case may be determined using techniques, such as a dose escalation study. The term "therapeutically effective amount" includes, for example, a prophylactically effective amount. An "effective amount" of a compound disclosed herein is an amount effective to achieve a desired pharmacologic effect or therapeutic improvement without undue adverse side effects. It is understood that "an effect amount" or "a therapeutically effective amount" can vary from subject to subject, due to variation in metabolism of Compound 1, age, weight, general condition of the subject, the condition being treated, the severity of the condition being treated, and the judgment of the prescribing physician. By way of example only, therapeutically effective amounts may be determined by routine experimentation, including but not limited to a dose escalation clinical trial.

The terms "enhance" or "enhancing" means to increase or prolong either in potency or duration a desired effect. By way of example, "enhancing" the effect of therapeutic agents refers to the ability to increase or prolong, either in potency or duration, the effect of therapeutic agents on during treatment of a disease, disorder or condition. An "enhancing-effective amount," as used herein, refers to an amount adequate to enhance the effect of a therapeutic agent in the treatment of a disease, disorder or condition. When used in a patient, amounts effective for this use will depend on the severity and course of the disease, disorder or condition, previous therapy, the patient's health status and response to the drugs, and the judgment of the treating physician.

The terms "inhibits", "inhibiting", or "inhibitor" of a kinase, as used herein, refer to inhibition of enzymatic phosphotransferase activity.

The term "irreversible inhibitor," as used herein, refers to a compound that, upon contact with a target protein (e.g., a kinase) causes the formation of a new covalent bond with or within the protein, whereby one or more of the target protein's biological activities (e.g., phosphotransferase activity) is diminished or abolished notwithstanding the subsequent presence or absence of the irreversible inhibitor.

The term "irreversible Btk inhibitor," as used herein,
forefers to an inhibitor of Btk that can form a covalent bond with an amino acid residue of Btk. In one embodiment, the irreversible inhibitor of Btk can form a covalent bond with a Cys residue of Btk; in particular embodiments, the irreversible inhibitor can form a covalent bond with a Cys 481 residue (or a homolog thereof) of Btk or a cysteine residue in the homologous corresponding position of another tyro-

The term "modulate," as used herein, means to interact with a target either directly or indirectly so as to alter the activity of the target, including, by way of example only, to enhance the activity of the target, to inhibit the activity of the target, to limit the activity of the target, or to extend the activity of the target.

As used herein, the term "modulator" refers to a compound that alters an activity of a molecule. For example, a modulator can cause an increase or decrease in the magnitude of a certain activity of a molecule compared to the magnitude of the activity in the absence of the modulator. In certain embodiments, a modulator is an inhibitor, which decreases the magnitude of one or more activities of a molecule. In certain embodiments, an inhibitor completely prevents one or more activities of a molecule. In certain embodiments, a modulator is an activator, which increases the magnitude of at least one activity of a molecule. In certain embodiments the presence of a modulator results in an activity that does not occur in the absence of the modulator.

The term "prophylactically effective amount," as used herein, refers that amount of a composition applied to a patient which will relieve to some extent one or more of the symptoms of a disease, condition or disorder being treated. In such prophylactic applications, such amounts may depend on the patient's state of health, weight, and the like. It is considered well within the skill of the art for one to determine such prophylactically effective amounts by routine experimentation, including, but not limited to, a dose 20 escalation clinical trial.

The term "individual," "subject" or "patient" as used herein, refers to an animal which is the object of treatment, observation or experiment. By way of example only, a subject may be, but is not limited to, a mammal including, 25 but not limited to, a human.

The term "wet granulation" as used herein, refers to the formation of granules using a granulation liquid (water, organic solvent, or a solution).

The term "dry granulation" as used herein, refers to the 30 formation of granules without using a granulation liquid (water, organic solvent, or a solution).

The term "high-load solid tablet formulation" as used herein, refers to a solid tablet formulation comprising at least 50% w/w of ibrutinib per tablet.

As used herein, the  $IC_{50}$  refers to an amount, concentration or dosage of a particular test compound that achieves a 50% inhibition of a maximal response, such as inhibition of Btk, in an assay that measures such response.

As used herein,  $\mathrm{EC}_{50}$  refers to a dosage, concentration or 40 amount of a particular test compound that elicits a dosedependent response at 50% of maximal expression of a particular response that is induced, provoked or potentiated by the particular test compound.

Pharmaceutical Compositions/Formulations

A pharmaceutical composition or pharmaceutical formulation, as used herein, refers to a mixture of Compound 1 with other chemical components, such as carriers, stabilizers, diluents, dispersing agents, suspending agents, thickening agents, and/or excipients. The pharmaceutical composition facilitates administration of the compound to a mammal. The compounds can be used singly or in combination with one or more therapeutic agents as components of mixtures.

The term "pharmaceutical combination" as used herein, 55 means a product that results from the mixing or combining of more than one active ingredient and includes both fixed and non-fixed combinations of the active ingredients. The term "fixed combination" means that the active ingredients, e.g. Compound 1 and a co-agent, are both administered to a patient simultaneously in the form of a single entity or dosage. The term "non-fixed combination" means that the active ingredients, e.g. Compound 1 and a co-agent, are administered to a patient as separate entities either simultaneously, concurrently or sequentially with no specific intervening time limits, wherein such administration provides effective levels of the two compounds in the body of the

patient. The latter also applies to cocktail therapy, e.g. the administration of three or more active ingredients.

In some embodiments, crystalline Compound 1 is incorporated into pharmaceutical compositions to provide solid oral dosage forms, such as powders, immediate release formulations, controlled release formulations, fast melt formulations, tablets, capsules, pills, delayed release formulations, extended release formulations, pulsatile release formulations, multiparticulate formulations, and mixed immediate and controlled release formulations.

Ibrutinib is currently used in the clinic at a unit dose of 420 mg or 560 mg which is administered orally in three or four capsules comprising 140 mg ibrutinib per capsule. High load tablet formulations would allow administration of one tablet per dose. However, high load tablet formulations that meet pharmaceutically acceptable properties, such as, suitable compressibility, compactibility, granulate flowability, granulate density, integrity during manufacture, shipping and storage, proper hardness, stability, swallowbility and disintegration properties when administered, are considerately more difficult to prepare than capsule formations due to the limited amount of excipients that can be used to adjust the tablet properties. Further, tablet formulations tend to have lower  $C_{max}$  as compared with the capsule formulations due to the process of its disintegration and absorption after administration, especially for ibrutinib which has a very low water solubility. It is challenging to prepare high load tablet formulations of ibrutinib that possess both pharmaceutically acceptable properties and desired PK properties, such as a high  $C_{max}$ .

In some embodiments is a pharmaceutical composition comprising ibrutinib, wherein ibrutinib is a compound with the structure of Compound 1,

and wherein the pharmaceutical composition comprises at least 50% w/w of ibrutinib.

In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises at least about 20% w/w of ibrutinib. In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises about 20% w/w to about 90% w/w of ibrutinib. In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises about 30% w/w to about 90% w/w of ibrutinib. In another embodiment is a pharmaceutical composition co

prising ibrutinib, wherein the pharmaceutical composition comprises about 40% w/w to about 90% w/w of ibrutinib. In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises about 50% w/w to about 90% w/w of ibrutinib. In 5 another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises about 40% w/w to about 80% w/w of ibrutinib. In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition 10 comprises about 50% w/w to about 80% w/w of ibrutinib. In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises about 60% w/w to about 80% w/w of ibrutinib. In another embodiment is a pharmaceutical composition com- 15 prising ibrutinib, wherein the pharmaceutical composition comprises about 50% w/w to about 75% w/w of ibrutinib. In another embodiment is a pharmaceutical composition comprising ibrutinib, wherein the pharmaceutical composition comprises about 60% w/w to about 75% w/w of ibrutinib. In 20 another embodiment is a pharmaceutical composition comprising at least 50% w/w of ibrutinib, wherein the pharmaceutical composition comprises intragranular and extragranular ingredients. In another embodiment is a pharmaceutical composition comprising at least 50% w/w of 25 ibrutinib, wherein the pharmaceutical composition is prepared using a wet granulation method. In another embodiment is a pharmaceutical composition comprising at least 50% w/w of ibrutinib, further comprising at least one pharmaceutically acceptable excipient.

In some embodiments, the pharmaceutical compositions described herein are prepared by a process comprising a wet granulation method.

In another embodiment is a solid tablet formulation comprising ibrutinib, wherein the solid tablet formulation 35 comprises at least about 20% w/w of ibrutinib. In another embodiment is a solid tablet formulation comprising ibrutinib, wherein the solid tablet formulation comprises about 20% w/w to about 90% w/w of ibrutinib. In another embodiment is a high-load solid tablet formulation comprising at 40 least 20% w/w or 30% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a high-load solid tablet formulation comprising at least 40% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a 45 high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a high-load solid tablet formulation comprising about 30% w/w to about 90% w/w of ibrutinib, and one or more pharmaceutically accept- 50 able excipients. In another embodiment is a high-load solid tablet formulation comprising about 40% w/w to about 90% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a high-load solid tablet formulation comprising about 50% w/w to about 90% 55 w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a high-load solid tablet formulation comprising about 40% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a high-load solid 60 tablet formulation comprising about 50% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a high-load solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a high-load solid tablet formulation comprising about 50% w/w to about 75%

32

w/w of ibrutinib, and one or more pharmaceutically acceptable excipients. In another embodiment is a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients.

In another embodiment is a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are present in an amount from about 10% w/w to about 50% w/w. In another embodiment is a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are present in an amount from about 10% w/w to about 50% w/w. In another embodiment is a highload solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are present in an amount from about 20% w/w to about 40% w/w. In another embodiment is a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are present in an amount from about 25% w/w to about 40% w/w.

In another embodiment is a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are selected from the group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants. In some embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose (e.g., Dipac®), dextrose, dextrates, maltodextrin, mannitol, xylitol (e.g., Xylitab®), sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose (e.g., Avicel®), microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is present in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrystalline cellulose. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent comprises lactose and microcrystalline cellulose. In some embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an amount of about 14% w/w and microcrystalline cellulose is present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose (e.g., Methocel®), croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked

carboxymethylcellulose, cross-linked croscarmellose, crosslinked starch such as sodium starch glycolate, cross-linked polymer such as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the disintegrating agent is croscarmellose 5 sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to about 10% w/w, about 5% w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one 10 excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone (e.g., PVP K15, PVP K19, PVP K25, PVP K30, Povidone® CL, Kollidon® CL, Polyplasdone® XL-10, and Povidone® K-12). In some embodiments, the polyvinylpyrrolidone is present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, about 0.1% w/w to about 20 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipient is a surfactant. In some embodiments, the surfactant is sodium lauryl sulfate (SLS). In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some embodiments, at least one 30 excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to about 1.5% w/w, 35 about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium stearate is present in 40 an amount from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose sodium, sodium lau- 45 ryl sulfate, colloidal silicon dioxide and magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose, microcrystalline 50 cellulose, croscarmellose sodium, and hydroxypropyl cellulose; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment, the intragranular excipients comprise:

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;

microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% 60 w/w, or about 8% w/w to about 15% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and

hydroxypropyl cellulose in an amount from about 0% 65 w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and

34

the extragranular excipients comprise:

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 5% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w;

colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment is a high-load solid tablet formu15 lation comprising at least 50% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose, microcrystalline
cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and
croscarmellose sodium; and the extragranular excipients
20 comprise croscarmellose sodium, sodium lauryl sulfate,
colloidal silicon dioxide, and magnesium stearate. In another
embodiment, the intragranular excipients comprise:

lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;

microcrystalline cellulose in an amount from about 1% w/w to about 10% w/w, about 2% w/w to about 5% w/w;

polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

croscarmellose sodium in an amount from about 1% w/w to about 10% w/w, or about 3% w/w to about 7% w/w; and

sodium lauryl sulfate in an amount from about 0% w/w to about 2% w/w, about 0.5% w/w to about 1.5% w/w;

the extragranular excipients comprise:

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w or about 0% w/w to about 4% w/w;

colloidal silicon dioxide in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w

In another embodiment is a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are selected from the group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants. In some embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, mannitol, xylitol, sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is present in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrystalline cellulose. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is

present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent comprises lactose and microcrystalline cellulose. In some embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an amount of about 14% w/w and  $_{15}$ microcrystalline cellulose is present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, a pregelatinized starch, a 20 sodium starch, methylcrystalline cellulose, methylcellulose, croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymethylcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, cross-linked polymer such 25 as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to about 10% w/w, about 5% 30 w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the polyvinylpyrrolidone is 35 present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, 40 about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipient is a surfactant. In some embodi- 45 ments, the surfactant is sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some 50 embodiments, at least one excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to 55 about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium 60 the extragranular excipients comprise stearate is present in an amount from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose 65 sodium, sodium lauryl sulfate, colloidal silicon dioxide and magnesium stearate.

36

In another embodiment is a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment, the intragranular excipients comprise:

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;

microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and

hydroxypropyl cellulose in an amount from about 0% w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and

the extragranular excipients comprise:

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 5% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w:

colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment is a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose. microcrystalline cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment, the intragranular excipients comprise:

lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;

microcrystalline cellulose in an amount from about 1% w/w to about 10% w/w, about 2% w/w to about 5%

polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

croscarmellose sodium in an amount from about 1% w/w to about 10% w/w, or about 3% w/w to about 7% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 2% w/w, about 0.5% w/w to about 1.5% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w or about 0% w/w to about 4% w/w;

colloidal silicon dioxide in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6%

In another embodiment is a high-load solid tablet formulation comprising about 50% w/w to about 80% w/w of 5 ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are selected from the group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants. In some embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, mannitol, xylitol, sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc. In some 15 embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is present in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and 20 lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrystalline cellulose. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 20% w/w, 25 about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellu- 30 lose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent comprises lactose and microcrystalline cellulose. In some embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and 35 microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an amount of about 14% w/w and microcrystalline cellulose is present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at 40 least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose, croscarmellose, croscarmellose sodium, cross-linked 45 the extragranular excipients comprise: sodium carboxymethylcellulose, cross-linked carboxymethvlcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, cross-linked polymer such as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the 50 disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to about 10% w/w, about 5% w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 55 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the polyvinylpyrrolidone is present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, 60 the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation 65 comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodi-

38

ments, at least one excipient is a surfactant. In some embodiments, the surfactant is sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some embodiments, at least one excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium stearate is present in an amount from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide and magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising about 50% w/w to about 80% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment the intragranular excipients comprise:

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;

microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and

hydroxypropyl cellulose in an amount from about 0% w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 5% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w:

colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment is a high-load solid tablet formulation comprising about 50% w/w to about 80% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, microcrystalline cellulose, colloidal silicon dioxide, and magnesium stearate. In another embodiment the intragranular excipients comprise:

lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;

polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

croscarmellose sodium in an amount from about 1% w/w 5 to about 10% w/w, or about 3% w/w to about 7% w/w; and

sodium lauryl sulfate in an amount from about 0% w/w to about 2% w/w, about 0.5% w/w to about 1.5% w/w; and

the extragranular excipients comprise:

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w or about 0% w/w to about 4% w/w; 15

microcrystalline cellulose in an amount from about 1% w/w to about 10% w/w, about 2% w/w to about 5% w/w;

colloidal silicon dioxide in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 20 0.6% w/w; and

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w

In another embodiment is a high-load solid tablet formu- 25 lation comprising about 60% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are selected from the group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants. In some 30 embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, mannitol, xylitol, sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, 35 microcrystalline cellulose, microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is present in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 40 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrystalline cellulose. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is 45 present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent 50 is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent comprises lactose and microcrystalline cellulose. In some embodiments, the lactose is pres- 55 ent in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an amount of about 14% w/w and microcrystalline cellulose is present in an amount from 60 about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose, 65 croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymeth40

ylcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, cross-linked polymer such as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to about 10% w/w, about 5% w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the polyvinylpyrrolidone is present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipient is a surfactant. In some embodiments, the surfactant is sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some embodiments, at least one excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium stearate is present in an amount from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide and magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment the intragranular excipients comprise

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;

microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and

hydroxypropyl cellulose in an amount from about 0% w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 5% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or 5 about 5% w/w to about 6% w/w;

colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.1% w/w 10 to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment is a high-load solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, and intragranular and extragranular excipients; 15 wherein the intragranular excipients comprise lactose, microcrystalline cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment the intragranular excipients comprise:

lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;

microcrystalline cellulose in an amount from about 1% 25 w/w to about 10% w/w, about 2% w/w to about 5% w/w;

polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

croscarmellose sodium in an amount from about 1% w/w 30 to about 10% w/w, or about 3% w/w to about 7% w/w; and

sodium lauryl sulfate in an amount from about 0% w/w to about 2% w/w, about 0.5% w/w to about 1.5% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w; sodium lauryl sulfate in an amount from about 0% w/w to

about 10% w/w or about 0% w/w to about 4% w/w; colloidal silicon dioxide in an amount from about 0.4% w/w to about 0.8% w/w or about 0.5% w/w to about 0.8% w/w or about 0.5% w/w to about

w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and magnesium stearate in an amount from about 0.4% w/w

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% 45 w/w.

In another embodiment is a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients wherein the one or more excipients are selected 50 from the group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants. In some embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, 55 mannitol, xylitol, sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is 60 present in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrys- 65 talline cellulose. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is

42

present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent comprises lactose and microcrystalline cellulose. In some embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an amount of about 14% w/w and microcrystalline cellulose is present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose, croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymethylcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, cross-linked polymer such as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to about 10% w/w, about 5% w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the polyvinylpyrrolidone is present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipient is a surfactant. In some embodiments, the surfactant is sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some embodiments, at least one excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium stearate is present in an amount from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide and magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment the intragranular excipients comprise:

- lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;
- microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w;
- croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and
- hydroxypropyl cellulose in an amount from about 0% 20 w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and

the extragranular excipients comprise:

- croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or 25 about 2% w/w to about 5% w/w;
- sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w:
- colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and
- magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment is a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the intragranular excipients comprise lactose, 40 microcrystalline cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment the intragranular 45 excipients comprise:

lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;

- microcrystalline cellulose in an amount from about 1% w/w to about 10% w/w, about 2% w/w to about 5% 50 w/w:
- polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;
- croscarmellose sodium in an amount from about 1% w/w to about 10% w/w, or about 3% w/w to about 7% w/w; 55 and
- sodium lauryl sulfate in an amount from about 0% w/w to about 2% w/w, about 0.5% w/w to about 1.5% w/w; and

60

the extragranular excipients comprise:

- croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;
- sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w or about 0% w/w to about 4% w/w;
- colloidal silicon dioxide in an amount from about 0.4% 65 w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

44

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w

In another embodiment is a high-load solid tablet formulation comprising:

- a) about 69% w/w to about 71% w/w of ibrutinib,
- b) about 13% w/w to about 15% w/w of lactose monohydrate,
- c) about 2% w/w to about 5% w/w of microcrystalline cellulose.
- d) about 1% w/w to about 3% w/w of polyvinylpyrrolidone.
- e) about 6% w/w to about 8% w/w of croscarmellose sodium.
- f) about 1% w/w to about 4% w/w of sodium lauryl sulfate.
- g) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
- h) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising:

- a) about 70% w/w of ibrutinib,
- b) about 14% w/w of lactose monohydrate,
- c) about 5% w/w of microcrystalline cellulose,
- d) about 2% w/w of polyvinylpyrrolidone,
- e) about 7% w/w of croscarmellose sodium,
- f) about 1% w/w of sodium lauryl sulfate,
- g) about 0.5% w/w of colloidal silicon dioxide, and
- h) about 0.5% w/w of magnesium stearate.

In one embodiment, the tablet formulation is as described above and croscarmellose sodium is about 5% intra and about 2% extra. In another embodiment, sodium lauryl sulfate is about 1% intra and about 0% extra.

In another embodiment is a high-load solid tablet formulation comprising:

- a) about 70% w/w of ibrutinib,
- b) about 14% w/w of lactose monohydrate,
- c) about 2% w/w of microcrystalline cellulose,
- d) about 2% w/w of polyvinylpyrrolidone,
- e) about 7% w/w of croscarmellose sodium,
- f) about 4% w/w of sodium lauryl sulfate,
- g) about 0.5% w/w of colloidal silicon dioxide, and
- h) about 0.5% w/w of magnesium stearate.

In one embodiment, the tablet formulation is as described above and croscarmellose sodium is about 5% intra and about 2% extra. In another embodiment, sodium lauryl sulfate is about 1% intra and about 3% extra.

In another embodiment, the high-load solid tablet comprises lactose, polyvinylpyrrolidone, sodium lauryl sulfate, crospovidone, colloidal silicon dioxide, and magnesium stearate. In another embodiment is a high-load solid tablet formulation comprising:

- a) about 65% w/w to about 75% w/w, or about 70% w/w of ibrutinib,
- b) about 14% w/w to about 18% w/w, or about 16% w/w of lactose monohydrate,
- c) about 1% w/w to about 3% w/w, or about 2% w/w of polyvinylpyrrolidone,
- d) about 0.5% w/w to about 1.5% w/w, or about 1% w/w of sodium lauryl sulfate,
- e) about 5% w/w to about 15% w/w, or about 10% w/w of crospovidone,
- f) about 0.3% w/w to about 0.7% w/w, or about 0.5% w/w of colloidal silicon dioxide, and
- g) about 0.3% w/w to about 0.7% w/w, or about 0.5% w/w of magnesium stearate.

In another embodiment is a high-load solid tablet formulation comprising:

- a) about 59% w/w to about 61% w/w of ibrutinib,
- b) about 13% w/w to about 15% w/w of lactose,
- c) about 13% w/w to about 15% w/w of microcrystalline 5 cellulose,
- d) about 4% w/w to about 6% w/w of croscarmellose sodium,
- e) about 5% w/w to about 7% w/w of sodium lauryl sulfate,
- f) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
- g) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In some embodiments, the total weight of a tablet is about 15 934 mg

In another embodiment is a high-load solid tablet formulation comprising:

- a) about 59% w/w to about 61% w/w of ibrutinib,
- b) about 13% w/w to about 14% w/w of lactose,
- c) about 13% w/w to about 14% w/w of microcrystalline cellulose,
- d) about 2% w/w to about 3% w/w of croscarmellose sodium(intragranular),
- e) about 0.8% w/w to about 1.2% w/w of hydroxypropyl 25 cellulose.
- f) about 2% w/w to about 3% w/w of croscarmellose sodium(extragranular),
- g) about 5.5 to about 6.5% w/w of sodium lauryl sulfate,
- h) about 0.4% w/w to about 0.6% w/w of colloidal silicon 30 dioxide, and
- i) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In some embodiments, the total weight of a tablet is about 934 mg.

In another embodiment is a high-load solid tablet formulation comprising:

- a) about 69% w/w to about 71% w/w of ibrutinib,
- b) about 8% w/w to about 9% w/w of lactose,
- c) about 8 to about 9% w/w of microcrystalline cellulose, 40
- d) about 2.5 to about 3.5% w/w of croscarmellose sodium (intragranular),
- e) about 2.5 to about 3.5% w/w of croscarmellose sodium (extragranular),
- g) about 5.5 to about 6.5% w/w of sodium lauryl sulfate,  $\,$  45
- h) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
- i) about 0.4% w/w to about 0.6% w/w of magnesium

In some embodiments of a tablet described herein the total 50 weight of the tablet is about 50 mg to about 1.2 g, such as about 50 mg, about 100 mg, about 200 mg, about 400 mg, about 600 mg, about 800 mg, or about 1.2 g, or any range between any two of the values, end points inclusive. In some embodiments, the total weight of a tablet is about 800 mg. 55

In some embodiments of the high-load solid tablet formulations described herein, the ibrutinib is in an amount of about 35 mg to about 840 mg per tablet, such as about 35 mg, about 70 mg, about 140 mg, about 280 mg, about 420 mg, about 560 mg, or about 840 mg, or any range between 60 any two of the values, end points inclusive. In some embodiments of the high-load solid tablet formulations described herein, the ibrutinib is in an amount of about 560 mg. In some embodiments of the high-load solid tablet formulations described herein, the ibrutinib is in micronized form. In 65 some embodiments of the high-load solid tablet formulations described herein, the formulation is used for once a day

46

dosing. In some embodiments of the high-load solid tablet formulations described herein, the formulation is in an oral dosage form containing a therapeutically effective amount of ibrutinib.

In some embodiments, the high-load solid tablet formulations described herein are prepared by a process comprising a wet granulation method.

In another embodiment is a method of treating a disease in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

In another embodiment is a method of treating an autoimmune disease in a patient in need of such treatment,
comprising administering to the patient a therapeutically
effective amount of a pharmaceutical composition or formulation described herein. In some embodiments, the autoimmune disease is rheumatoid arthritis or lupus. In another
embodiment is a method of treating rheumatoid arthritis in
a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a
pharmaceutical composition or formulation described
herein. In another embodiment is a method of treating lupus
in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a
pharmaceutical composition or formulation described
herein.

In another embodiment is a method of treating a heteroimmune disease in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein. In some embodiments, the cancer is a B-cell proliferative disorder. In some embodiments, the cancer is a B-cell proliferative disorder and the B-cell proliferative disorder is diffuse large B cell lymphoma, follicular lymphoma or chronic lymphocytic leukemia. In some embodiments, the cancer is a B-cell proliferative disorder and the B-cell proliferative disorder is diffuse large B cell lymphoma. In some embodiments, the cancer is a B-cell proliferative disorder and the B-cell proliferative disorder and the B-cell proliferative disorder is follicular lymphoma.

In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is a B cell malignancy. In some embodiments, the cancer is a B cell malignancy and the B cell malignancy selected from chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL), mantle cell lymphoma (MCL), diffuse large B Cell lymphoma (DLBCL), and multiple myeloma. In some embodiments, the cancer is a B cell malignancy and the B cell malignancy is chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL). In some embodiments, the cancer is a B cell malignancy and the B cell malignancy is mantle cell lymphoma (MCL). In some embodiments, the cancer is a B cell malignancy and the B cell malignancy is diffuse large B Cell lymphoma (DLBCL). In some embodiments, the cancer is a B cell malignancy and the B cell malignancy is multiple myeloma.

In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administer-

ing to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is a lymphoma, leukemia or a solid tumor. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising 5 administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is a lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering 10 to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is a leukemia. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a thera- 15 peutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is a solid tumor.

In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administer- 20 ing to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is diffuse large B cell lymphoma, follicular lymphoma, chronic lymphocytic lymphoma, chronic lymphocytic leukemia, B-cell prolymphocytic leu- 25 kemia, lymphoplasmacytic lymphoma/Waldenstrom macroglobulinemia, splenic marginal zone lymphoma, plasma cell myeloma, plasmacytoma, extranodal marginal zone B cell lymphoma, nodal marginal zone B cell lymphoma, mantle cell lymphoma, mediastinal (thymic) large B cell lym- 30 phoma, intravascular large B cell lymphoma, primary effusion lymphoma, burkitt lymphoma/leukemia, or lymphomatoid granulomatosis. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically 35 effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is diffuse large B cell lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically 40 effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is follicular lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective 45 amount of a pharmaceutical composition or formulation described herein, wherein the cancer is chronic lymphocytic lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective 50 amount of a pharmaceutical composition or formulation described herein, wherein the cancer is chronic lymphocytic leukemia. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective 55 amount of a pharmaceutical composition or formulation described herein, wherein the cancer is B-cell prolymphocytic leukemia. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically 60 effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is lymphoplasmacytic lymphoma/Waldenstrom macroglobulinemia. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering 65 to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein,

48

wherein the cancer is splenic marginal zone lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is plasma cell myeloma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is plasmacytoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is extranodal marginal zone B cell lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is nodal marginal zone B cell lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is mantle cell lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is mediastinal (thymic) large B cell lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is intravascular large B cell lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is primary effusion lymphoma. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is burkitt lymphoma/leukemia. In another embodiment is a method of treating cancer in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein, wherein the cancer is lymphomatoid granulomatosis.

In some embodiments, the composition is for use in treatment of a sarcoma or carcinoma. In some embodiments, the composition is for use in treatment of a sarcoma. In some embodiments, the composition is for use in treatment of a carcinoma. In some embodiments, the sarcoma is selected from alveolar rhabdomyosarcoma; alveolar soft part sarcoma; ameloblastoma; angiosarcoma; chondrosarcoma; chordoma; clear cell sarcoma of soft tissue; dedifferentiated liposarcoma; desmoid; desmoplastic small round cell tumor; embryonal rhabdomyosarcoma; epithelioid fibrosarcoma; epithelioid hemangioendothelioma; epithelioid sarcoma; esthesioneuroblastoma; Ewing sarcoma; extrarenal rhabdoid tumor; extraskeletal myxoid chondrosarcoma; extrasketetal osteosarcoma; fibrosarcoma; giant cell tumor; hemangiopericytoma; infantile fibrosarcoma; inflammatory myofibroblastic tumor; Kaposi sarcoma; leiomyosarcoma of bone; liposarcoma; liposarcoma of bone; malignant fibrous histiocytoma (MFH); malignant fibrous histiocytoma (MFH) of bone; malignant mesenchymoma; malignant peripheral nerve sheath tumor; mesenchymal chondrosarcoma; myxofibrosarcoma; myxoid liposarcoma; myxoinflammatory 5 fibroblastic sarcoma; neoplasms with perivascular epitheioid cell differentiation; osteosarcoma; parosteal osteosarcoma; neoplasm with perivascular epitheioid cell differentiation; periosteal osteosarcoma; pleomorphic liposarcoma; pleomorphic rhabdomyosarcoma; PNET/extraskeletal Ewing 10 tumor; rhabdomyosarcoma; round cell liposarcoma; small cell osteosarcoma; solitary fibrous tumor; synovial sarcoma; telangiectatic osteosarcoma. In some embodiments, the carcinoma is selected from an adenocarcinoma, squamous cell carcinoma, adenosquamous carcinoma, anaplastic carci- 15 noma, large cell carcinoma, or small cell carcinoma. In some embodiments, the solid tumor is selected from anal cancer; appendix cancer; bile duct cancer (i.e., cholangiocarcinoma); bladder cancer; brain tumor; breast cancer; HER2amplified breast cancer; cervical cancer; colon cancer; can- 20 cer of Unknown Primary (CUP); esophageal cancer; eye cancer; fallopian tube cancer; kidney cancer; renal cell carcinoma; liver cancer; lung cancer; medulloblastoma; melanoma; oral cancer; ovarian cancer; pancreatic cancer; pancreatic ductal cancer; parathyroid disease; penile cancer; 25 pituitary tumor; prostate cancer; rectal cancer; skin cancer; stomach cancer; testicular cancer; throat cancer; thyroid cancer; uterine cancer; vaginal cancer; or vulvar cancer. In some embodiments, the carcinoma is breast cancer. In some embodiments, the breast cancer is invasive ductal carci- 30 noma, ductal carcinoma in situ, invasive lobular carcinoma, or lobular carcinoma in situ. In some embodiments, the carcinoma is pancreatic cancer. In some embodiments, the pancreatic cancer is adenocarcinoma, or islet cell carcinoma. In some embodiments, the carcinoma is colorectal cancer. In 35 some embodiments, the colorectal cancer is adenocarcinoma. In some embodiments, the solid tumor is a colon polyp. In some embodiments, the colon polyp is associated with familial adenomatous polyposis. In some embodiments, the carcinoma is bladder cancer. In some embodi- 40 ments, the bladder cancer is transitional cell bladder cancer, squamous cell bladder cancer, or adenocarcinoma. In some embodiments, the carcinoma is lung cancer. In some embodiments, the lung cancer is a non-small cell lung cancer. In some embodiments, the non-small cell lung 45 cancer is adenocarcinoma, squamous-cell lung carcinoma, or large-cell lung carcinoma. In some embodiments, the non-small cell lung cancer is large cell lung cancer. In some embodiments, the lung cancer is a small cell lung cancer. In some embodiments, the carcinoma is prostate cancer. In 50 some embodiments, the prostate cancer is adenocarcinoma or small cell carcinoma. In some embodiments, the carcinoma is ovarian cancer. In some embodiments, the ovarian cancer is epithelial ovarian cancer. In some embodiments, the carcinoma is bile duct cancer. In some embodiments, the 55 bile duct cancer is proximal bile duct carcinoma or distal bile

In another embodiment is a method of treating mastocytosis in a patient in need of such treatment, comprising administering to the patient a therapeutically effective 60 amount of a pharmaceutical composition or formulation described herein.

In another embodiment is a method of treating osteoporosis or bone resorption disorders in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein. In another embodiment is a

method of treating osteoporosis in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein. In another embodiment is a method of treating bone resorption disorders in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

In another embodiment is a method of treating an inflammatory disease or condition in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

In another embodiment is a method of treating lupus in a patient in need of such treatment, comprising administering to the patient a therapeutically effective amount of a pharmaceutical composition or formulation described herein.

In another aspect is a process for preparing a pharmaceutical composition described herein wherein the process comprises a wet granulation method.

In another aspect is a process for preparing a pharmaceutical composition comprising ibrutinib, wherein ibrutinib is a compound with the structure of Compound 1,

the process comprises a wet granulation method; and the pharmaceutical composition comprises at least 50% w/w of ibrutinib

In another embodiment is a process for preparing a pharmaceutical composition comprising ibrutinib, wherein the process comprises a wet granulation method and the pharmaceutical composition comprises about 30% w/w to about 90% w/w of ibrutinib. In another embodiment is a process for preparing a pharmaceutical composition comprising ibrutinib, wherein the process comprises a wet granulation method and the pharmaceutical composition comprises about 40% w/w to about 90% w/w of ibrutinib. In another embodiment is a process for preparing a pharmaceutical composition comprising ibrutinib, wherein the process comprises a wet granulation method and the pharmaceutical composition comprises about 50% w/w to about 90% w/w of ibrutinib. In another embodiment is a process for preparing a pharmaceutical composition comprising ibrutinib, wherein the process comprises a wet granulation method and the pharmaceutical composition comprises about 40% w/w to about 80% w/w of ibrutinib. In another embodiment is a process for preparing a pharmaceutical

composition comprising ibrutinib, wherein the process comprises a wet granulation method and the pharmaceutical composition comprises about 50% w/w to about 80% w/w of ibrutinib. In another embodiment is a process for preparing a pharmaceutical composition comprising ibrutinib, wherein 5 the process comprises a wet granulation method and the pharmaceutical composition comprises about 60% w/w to about 80% w/w of ibrutinib. In another embodiment is a process for preparing a pharmaceutical composition comprising ibrutinib, wherein the process comprises a wet 10 granulation method and the pharmaceutical composition comprises about 50% w/w to about 75% w/w of ibrutinib. In another embodiment is a process for preparing a pharmaceutical composition comprising ibrutinib, wherein the process comprises a wet granulation method and the pharma- 15 ceutical composition comprises about 60% w/w to about 75% w/w of ibrutinib.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, about 50% w/w to about 90% w/w of 20 ibrutinib, about 50% w/w to about 80% w/w of ibrutinib, about 60% w/w to about 80% w/w of ibrutinib, or about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients, wherein the process comprises a wet granulation method. In another embodiment 25 provided is a process for preparing a high-load solid tablet formulation comprising at least 30% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a 30 high-load solid tablet formulation comprising at least 40% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation compris- 35 ing at least 50% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 30% w/w to about 90% w/w 40 of ibrutinib, and one or more pharmaceutically acceptable excipients, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 40% w/w to about 90% w/w of ibrutinib, and one or 45 more pharmaceutically acceptable excipients, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, and one or more pharmaceutically 50 acceptable excipients, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 40% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients, 55 wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients, wherein the process 60 comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients, wherein the process comprises a wet granulation 65 method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising

**52** 

about 50% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients, wherein the process comprises a wet granulation method.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, about 50% w/w to about 90% w/w of ibrutinib, about 50% w/w to about 80% w/w of ibrutinib, about 60% w/w to about 80% w/w of ibrutinib, or about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients present in a total amount from about 10% w/w to about 50% w/w, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients present in a total amount no more than about 50% w/w, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients present in a total amount from about 10% w/w to about 50% w/w, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients present in a total amount from about 20% w/w to about 50% w/w, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients present in a total amount from about 20% w/w to about 40% w/w, wherein the process comprises a wet granulation method. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients present in a total amount from about 25% w/w to about 40% w/w, wherein the process comprises a wet granulation method.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients selected from the group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants, wherein the process comprises a wet granulation method. In some embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, mannitol, xylitol, sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is present in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrystalline cellulose. In some embodiments, the diluent is microcrystalline cellulose and the

microcrystalline cellulose is present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In 5 some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent comprises lactose and microcrystalline cellulose. In some embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an 15 amount of about 14% w/w and microcrystalline cellulose is present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, 20 a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose, croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymethylcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, 25 cross-linked polymer such as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to 30 about 10% w/w, about 5% w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the 35 polyvinylpyrrolidone is present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% 40 w/w, about 0 to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipi- 45 ent is a surfactant. In some embodiments, the surfactant is sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to 50 about 6% w/w. In some embodiments, at least one excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 55 to about 5% w/w, 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium 60 stearate and the magnesium stearate is present in an amount from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvi- 65 nylpyrrolidone, croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide and magnesium stearate.

54

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the process comprises a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose, and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, wherein the process comprises a wet granulation method, the intragranular excipients comprise

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;

microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and

hydroxypropyl cellulose in an amount from about 0% w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 5% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w;

colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the process comprises a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium, and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising at least 50% w/w of ibrutinib, wherein the process comprises a wet granulation method, the intragranular excipients comprise

lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;

microcrystalline cellulose in an amount from about 1% w/w to about 10% w/w, about 2% w/w to about 5% w/w;

polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

croscarmellose sodium in an amount from about 1% w/w to about 10% w/w, or about 3% w/w to about 7% w/w; and

sodium lauryl sulfate in an amount from about 0% w/w to about 2% w/w, about 0.5% w/w to about 1.5% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w; sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w or about 0% w/w to about 4% w/w; colloidal silicon dioxide in an amount from about 0.4% 5 w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients selected from the group consisting of diluents, binders, disintegrating agents, 15 lubricants, glidants, and surfactants. In some embodiments, at least one excipient is a diluent, wherein the process comprises a wet granulation method. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, mannitol, xvlitol, 20 sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is present in an amount from 25 about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrystalline cellulose. In 30 some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, 5% w/w to about 20% w/w, about 8% w/w 35 to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent com- 40 prises lactose and microcrystalline cellulose. In some embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an 45 amount of about 14% w/w and microcrystalline cellulose is present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, 50 a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose, croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymethylcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, 55 cross-linked polymer such as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to 60 about 10% w/w, about 5% w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the polyvinylpyrrolidone is present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2%

56

w/w. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipient is a surfactant. In some embodiments, the surfactant is sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some embodiments, at least one excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium stearate is present in an amount from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5%w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide and magnesium stearate.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the process comprises a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose, and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, wherein the process comprises a wet granulation method, the intragranular excipients comprise

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;

microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and

hydroxypropyl cellulose in an amount from about 0% w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 5% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w;

colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the process comprises a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium, and 10 the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 90% w/w of ibrutinib, 15 wherein the process comprises a wet granulation method, the intragranular excipients comprise

lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;

microcrystalline cellulose in an amount from about 1% 20 w/w to about 10% w/w, about 2% w/w to about 5% w/w;

polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

croscarmellose sodium in an amount from about 1% w/w 25 to about 10% w/w, or about 3% w/w to about 7% w/w; and

sodium lauryl sulfate in an amount from about 0% w/w to about 2% w/w, about 0.5% w/w to about 1.5% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w; sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w or about 0% w/w to about 4% w/w; 35 colloidal silicon dioxide in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% 40 w/w.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients selected from the 45 group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants, wherein the process comprises a wet granulation method. In some embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, 50 sucrose, dextrose, dextrates, maltodextrin, mannitol, xylitol, sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the 55 diluent is lactose; and lactose is present in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some 60 embodiments, the diluent is microcrystalline cellulose. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15%

58

w/w. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent comprises lactose and microcrystalline cellulose. In some embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an amount of about 14% w/w and microcrystalline cellulose is present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose, croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymethylcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, cross-linked polymer such as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to about 10% w/w, about 5% w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the polyvinylpyrrolidone is present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipient is a surfactant. In some embodiments, the surfactant is sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some embodiments, at least one excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium stearate is present in an amount from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide and magnesium stearate.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 80% w/w of ibrutinib and intragranular and extragranular excipients, wherein the process comprises a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose

sodium, and hydroxypropyl cellulose, and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% by w/w to about 80% w/w of ibrutinib, wherein the process comprises a wet granulation method, the intragranular excipients comprise

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;

microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and

hydroxypropyl cellulose in an amount from about 0% w/w to about 2% w/w, about 0.1% w/w to about 1.1% 20 w/w, or about 0.1% w/w to about 1% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 5% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w;

colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 80% w/w of ibrutinib, and intragranular and extragranular excipients, wherein the process comprises a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium, and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment provided is a 45 process for preparing a high-load solid tablet formulation comprising about 50% w/w to about 80% w/w of ibrutinib, wherein the process comprises a wet granulation method, the intragranular excipients comprise

lactose in an amount from about 10% w/w to about 20% 50 w/w, or about 12% w/w to about 15% w/w;

microcrystalline cellulose in an amount from about 1% w/w to about 10% w/w, about 2% w/w to about 5% w/w:

polyvinylpyrrolidone in an amount from about 0% w/w to 55 about 5% w/w, about 1% w/w to about 3% w/w;

croscarmellose sodium in an amount from about 1% w/w to about 10% w/w, or about 3% w/w to about 7% w/w; and

sodium lauryl sulfate in an amount from about 0% w/w to 60 about 2% w/w, about 0.5% w/w to about 1.5% w/w;

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w; sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w or about 0% w/w to about 4% w/w;

60

colloidal silicon dioxide in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients selected from the group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants, wherein the process comprises a wet granulation method. In some embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, mannitol, xylitol, sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is present in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrystalline cellulose. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent comprises lactose and microcrystalline cellulose. In some embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an amount of about 14% w/w and microcrystalline cellulose is present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose, croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymethylcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, cross-linked polymer such as crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to about 10% w/w, about 5% w/w to about 10% w/w, about 6% w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the polyvinylpyrrolidone is present in an amount from about 0 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, about 0.1% w/w to about

1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipient is a surfactant. In some embodiments, the surfactant is 5 sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some embodiments, at least one excipient is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to about 1.5% w/w, about 0.4% 15 w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium stearate is present in an amount 20 from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose sodium, sodium lauryl sul- 25 fate, colloidal silicon dioxide and magnesium stearate.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, and intragranular and extragranular excipients, wherein the process comprises 30 a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose, and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In 35 another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, wherein the process comprises a wet granulation method, the intragranular excipients comprise

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w;

microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% 45 w/w, or about 8% w/w to about 15% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and

hydroxypropyl cellulose in an amount from about 0% 50 w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or 55 about 2% w/w to about 5% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w;

colloidal silicon dioxide in an amount from about 0.1% 60 w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about

60% w/w to about 80% w/w of ibrutinib, and intragranular and extragranular excipients, wherein the process comprises a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium, and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 80% w/w of ibrutinib, wherein the process comprises a wet granulation method, the intragranular excipients comprise

62

lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;

microcrystalline cellulose in an amount from about 1% w/w to about 10% w/w, about 2% w/w to about 5% w/w.

polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

croscarmellose sodium in an amount from about 1% w/w to about 10% w/w, or about 3% w/w to about 7% w/w; and

sodium lauryl sulfate in an amount from about 0% w/w to about 2% w/w, about 0.5% w/w to about 1.5% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w or about 0% w/w to about 4% w/w;

colloidal silicon dioxide in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and one or more pharmaceutically acceptable excipients selected from the group consisting of diluents, binders, disintegrating agents, lubricants, glidants, and surfactants, wherein the process comprises a wet granulation method. In some embodiments, at least one excipient is a diluent. In some embodiments, the diluent is selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, mannitol, xylitol, sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc. In some embodiments, the diluent is cellulose. In some embodiments, the diluent is the diluent is lactose; and lactose is present in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is lactose; and lactose is present in an amount of about 8.5% w/w or about 14% w/w. In some embodiments, the diluent is microcrystalline cellulose. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 20% w/w, about 1% w/w to about 10% w/w, about 1% w/w to about 5% w/w, 1% w/w to about 2% w/w, about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w. In some embodiments, the diluent is microcrystalline cellulose and the microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w or about 8.5% w/w or about 14% w/w. In some embodiments, the diluent comprises lactose and microcrystalline cellulose. In some

embodiments, the lactose is present in an amount of about 10% w/w to about 15% w/w and microcrystalline cellulose is present in an amount from about 1% w/w to about 6% w/w. In some embodiments, the lactose is present in an amount of about 14% w/w and microcrystalline cellulose is 5 present in an amount from about 2% w/w to about 5% w/w. In some embodiments, at least one excipient is a disintegrating agent. In some embodiments, the disintegrating agent is selected from the group consisting of natural starch, a pregelatinized starch, a sodium starch, methylcrystalline 10 cellulose, methylcellulose, croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymethylcellulose, cross-linked croscarmellose, cross-linked starch such as sodium starch glycolate, cross-linked polymer such as crospovidone, cross-linked 15 polyvinylpyrrolidone, sodium alginate, a clay, and a gum. In some embodiments, the disintegrating agent is croscarmellose sodium; and croscarmellose sodium is present in an amount from about 0 to about 20% w/w, about 1% w/w to about 10% w/w, about 5% w/w to about 10% w/w, about 6% 20 w/w to about 8% w/w, about 4% w/w to about 6% w/w, or about 2% w/w to about 4% w/w. In some embodiments, at least one excipient is a binder. In some embodiments, the binder is polyvinylpyrrolidone. In some embodiments, the polyvinylpyrrolidone is present in an amount from about 0 25 to about 10% w/w, about 1 to about 5% w/w, or about 2% w/w. In some embodiments, the binder is hydroxypropyl cellulose; and hydroxypropyl cellulose is present in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 0 to about 2% w/w, about 0.1% w/w to about 30 1.1% w/w, or about 0.1% w/w to about 1% w/w. In some embodiments, the formulation comprises lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose. In some embodiments, at least one excipient is a surfactant. In some embodiments, the surfactant is 35 sodium lauryl sulfate. In some embodiments, the surfactant is sodium lauryl sulfate in an amount from about 0 to about 10% w/w, about 0.5 to about 5% w/w, about 1 to about 4% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w. In some embodiments, at least one excipient 40 is a glidant. In some embodiments, the glidant is silica (colloidal silicon dioxide). In some embodiments, the glidant is silica (colloidal silicon dioxide) and the silica (colloidal silicon dioxide) is present in an amount from about 0 to about 5% w/w, 0.1% w/w to about 1.5% w/w, about 0.4% 45 w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, at least one excipient is a lubricant. In some embodiments, the lubricant is magnesium stearate. In some embodiments, the lubricant is magnesium stearate and the magnesium stearate is present in an amount 50 from about 0.01% w/w to about 5% w/w, 0.01% w/w to about 2% w/w, 0.1% w/w to about 0.7% w/w, or about 0.5% w/w to about 0.6% w/w. In some embodiments, the excipients comprise lactose, microcrystalline cellulose, polyvinylpyrrolidone, croscarmellose sodium, sodium lauryl sul- 55 fate, colloidal silicon dioxide and magnesium stearate.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the process comprises 60 a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose; and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In 65 another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60%

64

w/w to about 75% w/w of ibrutinib, wherein the process comprises a wet granulation method, the intragranular excipients comprise

lactose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 15% w/w, or about 8% w/w to about 14% w/w:

microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, or about 8% w/w to about 15% w/w;

croscarmellose sodium in an amount from about 0% w/w to about 10% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 4% w/w; and

hydroxypropyl cellulose in an amount from about 0% w/w to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 2% w/w to about 5% w/w, or about 2% w/w to about 5% w/w;

sodium lauryl sulfate in an amount from about 0% w/w to about 10% w/w, about 4% w/w to about 8% w/w, or about 5% w/w to about 6% w/w;

colloidal silicon dioxide in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, and intragranular and extragranular excipients; wherein the process comprises a wet granulation method, the intragranular excipients comprise lactose, microcrystalline cellulose, sodium lauryl sulfate, polyvinylpyrrolidone and croscarmellose sodium, and the extragranular excipients comprise croscarmellose sodium, sodium lauryl sulfate, colloidal silicon dioxide, and magnesium stearate. In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising about 60% w/w to about 75% w/w of ibrutinib, wherein the process comprises a wet granulation method, the intragranular excipients comprise

lactose in an amount from about 10% w/w to about 20% w/w, or about 12% w/w to about 15% w/w;

microcrystalline cellulose in an amount from about 1% w/w to about 10% w/w, about 2% w/w to about 5% w/w.

polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

croscarmellose sodium in an amount from about 1% w/w to about 10% w/w, or about 3% w/w to about 7% w/w; and

sodium lauryl sulfate in an amount from about 0% w/w to about 2% w/w, about 0.5% w/w to about 1.5% w/w; and

the extragranular excipients comprise

croscarmellose sodium in an amount from about 0% w/w to about 5% w/w, about 1% w/w to about 3% w/w;

sodium lauryl sulfate in an amount from about 9% w/w to about 10% w/w or about 0% w/w to about 4% w/w;

colloidal silicon dioxide in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w; and

magnesium stearate in an amount from about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.

65

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising ibrutinib, wherein the process comprises a wet granulation method, and the formulation comprises:

- a) about 69% w/w to about 71% w/w of ibrutinib.
- b) about 13% w/w to about 15% w/w of lactose,
- c) about 2% w/w to about 5% w/w of microcrystalline cellulose.
- d) about 1% w/w to about 3% w/w of polyvinylpyrrolidone
- e) about 6% w/w to about 8% w/w of croscarmellose sodium.
- f) about 1% w/w to about 4% w/w of sodium lauryl sulfate,
- g) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
- h) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising ibrutinib, wherein the process comprises a wet granulation method, and the formulation comprises:

- a) about 70% w/w of ibrutinib,
- b) about 14% w/w of lactose,
- c) about 5% w/w of microcrystalline cellulose,
- d) about 2% w/w of polyvinylpyrrolidone,
- e) about 7% w/w of croscarmellose sodium,
- f) about 1% w/w of sodium lauryl sulfate,
- g) about 0.5% w/w of colloidal silicon dioxide, and
- h) about 0.5% w/w of magnesium stearate.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising ibrutinib, wherein the process comprises a wet granulation method, and the formulation comprises:

- a) about 70% w/w of ibrutinib,
- b) about 14% w/w of lactose,
- c) about 2% w/w of microcrystalline cellulose,
- d) about 2% w/w of polyvinylpyrrolidone,
- e) about 7% w/w of croscarmellose sodium,
- f) about 4% w/w of sodium lauryl sulfate,
- g) about 0.5% w/w of colloidal silicon dioxide, and
- h) about 0.5% w/w of magnesium stearate.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising ibru- 45 tinib, wherein the process comprises a wet granulation method, and wherein the formulation comprises:

- a) about 65% w/w to about 75% w/w, or about 70% w/w of ibrutinib,
- b) about 14% w/w to about 18% w/w, or about 16% w/w 50 of lactose monohydrate,
- c) about 1% w/w to about 3% w/w, or about 2% w/w of polyvinylpyrrolidone,
- d) about 0.5% w/w to about 1.5% w/w, or about 1% w/w of sodium lauryl sulfate,
- e) about 5% w/w to about 15% w/w, or about 10% w/w of crospovidone,
- f) about 0.3% w/w to about 0.7% w/w, or about 0.5% w/w of colloidal silicon dioxide, and
- g) about 0.3% w/w to about 0.7% w/w, or about 0.5% w/w  $\,$  60 of magnesium stearate.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising ibrutinib, wherein the process comprises a wet granulation method, and wherein the formulation comprises:

- a) about 59% w/w to about 61% w/w of ibrutinib,
- b) about 13% w/w to about 15% w/w of lactose,

66

- c) about 13% w/w to about 15% w/w of microcrystalline cellulose.
- d) about 4% w/w to about 6% w/w of croscarmellose sodium.
- e) about 5% w/w to about 7% w/w of sodium lauryl sulfate.
- f) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
- g) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In some embodiments, the total weight of a tablet is about 934 mg.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising ibrutinib, wherein the process comprises a wet granulation method, and wherein the formulation comprises:

- a) about 59% w/w to about 61% w/w of ibrutinib,
- b) about 13% w/w to about 14% w/w of lactose,
- c) about 13% w/w to about 14% w/w of microcrystalline cellulose.
- d) about 2% w/w to about 3% w/w of croscarmellose sodium(intragranular),
- e) about 0.8% w/w to about 1.2% w/w of hydroxypropyl cellulose,
- f) about 2% w/w to about 3% w/w of croscarmellose sodium(extragranular),
- g) about 5.5 to about 6.5% w/w of sodium lauryl sulfate,
- h) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
- i) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In some embodiments, the total weight of a tablet is about 934 mg.

In another embodiment provided is a process for preparing a high-load solid tablet formulation comprising ibrutinib, wherein the process comprises a wet granulation method, and wherein the formulation comprises:

- a) about 69% w/w to about 71% w/w of ibrutinib,
- b) about 8% w/w to about 9% w/w of lactose,
  - c) about 8 to about 9% w/w of microcrystalline cellulose,
  - d) about 2.5 to about 3.5% w/w of croscarmellose sodium (intragranular),
  - e) about 2.5 to about 3.5% w/w of croscarmellose sodium (extragranular),
  - g) about 5.5 to about 6.5% w/w of sodium lauryl sulfate,
  - h) about 0.4% w/w to about 0.6% w/w of colloidal silicon dioxide, and
  - i) about 0.4% w/w to about 0.6% w/w of magnesium stearate.

In some embodiments, the total weight of a tablet is about 800 mg.

In some embodiments of the high-load solid tablet formulations described herein comprising ibrutinib and prepared using a wet granulation method, the ibrutinib is in an amount of about 560 mg. In some embodiments of the high-load solid tablet formulations described herein comprising ibrutinib and prepared using a wet granulation method, the ibrutinib is in micronized form. In some embodiments of the high-load solid tablet formulations described herein comprising ibrutinib and prepared using a wet granulation method, the formulation is used for once a day dosing. In some embodiments of the high-load solid tablet formulations described herein comprising ibrutinib and prepared using a wet granulation method, the formulation is in an oral dosage form containing a therapeutically effective amount of ibrutinib.

Moreover, the pharmaceutical compositions described herein, which include Compound 1 can be formulated into any suitable dosage form, including but not limited to, solid oral dosage forms, controlled release formulations, fast melt formulations, effervescent formulations, tablets, powders, pills, capsules, delayed release formulations, extended release formulations, pulsatile release formulations, multiparticulate formulations, and mixed immediate release and controlled release formulations. In some embodiments, the tablets described herein are for immediate release and do not comprise a viscosity increasing agent, such as poloxamer or glyceryl behenate.

In some embodiments, the solid dosage forms disclosed herein may be in the form of a tablet, including a suspension tablet, a fast-melt tablet, a bite-disintegration tablet, a rapid-disintegration tablet, an effervescent tablet, or a caplet. In other embodiments, the pharmaceutical formulation is in the form of a powder. In still other embodiments, the pharmaceutical formulation is in the form of a tablet, including but not limited to, a fast-melt tablet. Additionally, pharmaceutical formulations described herein may be administered as a single capsule or in multiple capsule dosage form. In some embodiments, the pharmaceutical formulation is administered in two, or three, or four, tablets.

In some embodiments, the compositions described herein are prepared by mixing particles of Compound 1 with one or more pharmaceutical excipients to form a bulk blend composition. When referring to these bulk blend compositions as homogeneous, it is meant that the particles of Compound 1 are dispersed evenly throughout the composition so that the composition may be readily subdivided into equally effective unit dosage forms, such as tablets, pills, and capsules. The individual unit dosages may also include film coatings, which disintegrate upon oral ingestion or upon contact with diluent.

In some embodiments, the wet granulation method comprises granulating the mixture of ibrutinib and the intragranular excipients with a granulation liquid, such as purified water, under granulation conditions, such as high shear granulation conditions, to form granules.

In some embodiments, the compositions or formulations described herein are prepared by a method comprising (1) mixing ibrutinib with the intragranular excipients such as 45 filler, binder, disintegrant and surfactant; (2) granulating the mixture of ibrutinib and the intragranular excipients with purified water or an aqueous binder solution under high shear granulation conditions to form granules; (3) drying the granules to form dried granules; (4) milling the dried granules; (5) blending the milled granules with the extragranular excipients such as filler, disintegrant, surfactant and lubricant; and (6) compressing the mixture of milled granules and the extragranular excipients to form tablets.

The pharmaceutical compositions or formulations described herein can further include a flavoring agent, sweetening agent, colorant, antioxidant, preservative, or one or more combination thereof. In still other aspects, using standard coating procedures, such as those described in 60 *Remington's Pharmaceutical Sciences*, 20th Edition (2000), a film coating is provided around the formulation of Compound 1. In one embodiment, some or all of the particles of the Compound 1 are coated. In another embodiment, some or all of the particles of the Compound 1 are microencapsulated. In still another embodiment, the particles of the Compound 1 are not microencapsulated and are uncoated.

68

Suitable antioxidants for use in the compositions or formulations described herein include, for example, e.g., butylated hydroxytoluene (BHT), sodium ascorbate, and tocopherol.

It should be appreciated that there is considerable overlap between additives used in the solid dosage forms described herein. Thus, the above-listed additives should be taken as merely exemplary, and not limiting, of the types of additives that can be included in the compositions or formulations described herein. The amounts of such additives can be readily determined by one skilled in the art, according to the particular properties desired.

Compressed tablets are solid dosage forms prepared by compacting the bulk blend of the formulations described above. In various embodiments, compressed tablets which are designed to dissolve in the mouth will include one or more flavoring agents. In other embodiments, the compressed tablets will include a film surrounding the final compressed tablet. In some embodiments, the film coating can provide a delayed release of Compound 1 from the formulation. In other embodiments, the film coating aids in patient compliance (e.g., Opadry® coatings or sugar coating). Film coatings including Opadry® typically range from about 1% to about 3% of the tablet weight. In other embodiments, the compressed tablets include one or more excipients.

In some embodiments, the compositions or formulations described herein can be formulated as enteric coated delayed release oral dosage forms, i.e., as an oral dosage form of a pharmaceutical composition as described herein which utilizes an enteric coating to affect release in the small intestine of the gastrointestinal tract. The enteric coated dosage form may be a compressed or molded or extruded tablet/mold (coated or uncoated) containing granules, powder, pellets, beads or particles of the active ingredient and/or other composition components, which are themselves coated or uncoated. The enteric coated oral dosage form may also be a capsule (coated or uncoated) containing pellets, beads or granules of the solid carrier or the composition, which are themselves coated or uncoated.

The term "delayed release" as used herein refers to the delivery so that the release can be accomplished at some generally predictable location in the intestinal tract more distal to that which would have been accomplished if there had been no delayed release alterations. In some embodiments the method for delay of release is coating. Any coatings should be applied to a sufficient thickness such that the entire coating does not dissolve in the gastrointestinal fluids at pH below about 5, but does dissolve at pH about 5 and above. It is expected that any anionic polymer exhibiting a pH-dependent solubility profile can be used as an enteric coating in the methods and compositions described herein to achieve delivery to the lower gastrointestinal tract. In some embodiments the polymers described herein are anionic carboxylic polymers. In other embodiments, the polymers and compatible mixtures thereof, and some of their properties, include, but are not limited to:

Shellac, also called purified lac, a refined product obtained from the resinous secretion of an insect. This coating dissolves in media of pH >7;

Acrylic polymers. The performance of acrylic polymers (primarily their solubility in biological fluids) can vary based on the degree and type of substitution. Examples of suitable acrylic polymers include methacrylic acid copolymers and ammonium methacrylate copolymers. The Eudragit series E, L, S, RL, RS and NE (Rohm Pharma) are available as solubilized in organic solvent, aqueous disper-

0~ 10,010,007 =

sion, or dry powders. The Eudragit series RL, NE, and RS are insoluble in the gastrointestinal tract but are permeable and are used primarily for colonic targeting. The Eudragit series E dissolve in the stomach. The Eudragit series L, L-30D and S are insoluble in stomach and dissolve in the 5 intestine;

69

Cellulose Derivatives. Examples of suitable cellulose derivatives are: ethyl cellulose; reaction mixtures of partial acetate esters of cellulose with phthalic anhydride. The performance can vary based on the degree and type of substitution. Cellulose acetate phthalate (CAP) dissolves in pH >6. Aquateric (FMC) is an aqueous based system and is a spray dried CAP psuedolatex with particles <1 m. Other components in Aquateric can include pluronics, Tweens, and acetylated monoglycerides. Other suitable cellulose deriva- 15 tives include: cellulose acetate trimellitate (Eastman); methylcellulose (Pharmacoat, Methocel); hydroxypropylmethyl cellulose phthalate (HPMCP); hydroxypropylmethyl cellulose succinate (HPMCS); and hydroxypropylmethylcellulose acetate succinate (e.g., AQOAT (Shin Etsu)). The 20 performance can vary based on the degree and type of substitution. For example, HPMCP such as, HP-50, HP-55, HP-55S, HP-55F grades are suitable. The performance can vary based on the degree and type of substitution. For example, suitable grades of hydroxypropylmethylcellulose 25 acetate succinate include, but are not limited to, AS-LG (LF), which dissolves at pH 5, AS-MG (MF), which dissolves at pH 5.5, and AS-HG (HF), which dissolves at higher pH. These polymers are offered as granules, or as fine powders for aqueous dispersions; Poly Vinyl Acetate Phtha- 30 late (PVAP). PVAP dissolves in pH >5, and it is much less permeable to water vapor and gastric fluids.

In some embodiments, the coating can, and usually does, contain a plasticizer and possibly other coating excipients such as colorants, talc, and/or magnesium stearate, which are 35 well known in the art. Suitable plasticizers include triethyl citrate (Citroflex 2), triacetin (glyceryl triacetate), acetyl triethyl citrate (Citroflec A2), Carbowax 400 (polyethylene glycol 400), diethyl phthalate, tributyl citrate, acetylated monoglycerides, glycerol, fatty acid esters, propylene gly- 40 col, and dibutyl phthalate. In particular, anionic carboxylic acrylic polymers usually will contain 10-25% by weight of a plasticizer, especially dibutyl phthalate, polyethylene glycol, triethyl citrate and triacetin. Conventional coating techniques such as spray or pan coating are employed to apply 45 coatings. The coating thickness must be sufficient to ensure that the oral dosage form remains intact until the desired site of topical delivery in the intestinal tract is reached.

Colorants, detackifiers, surfactants, antifoaming agents, lubricants (e.g., camuba wax or PEG) may be added to the 50 coatings besides plasticizers to solubilize or disperse the coating material, and to improve coating performance and the coated product.

In other embodiments, the formulations described herein, which include Compound 1, are delivered using a pulsatile 55 dosage form. A pulsatile dosage form is capable of providing one or more immediate release pulses at predetermined time points after a controlled lag time or at specific sites. Many other types of controlled release systems known to those of ordinary skill in the art and are suitable for use with the 60 formulations described herein. Examples of such delivery systems include, e.g., polymer-based systems, such as polylactic and polyglycolic acid, plyanhydrides and polycaprolactone; porous matrices, nonpolymer-based systems that are lipids, including sterols, such as cholesterol, cholesterol esters and fatty acids, or neutral fats, such as mono-, di- and triglycerides; hydrogel release systems; silastic systems;

70

peptide-based systems; wax coatings, bioerodible dosage forms, compressed tablets using conventional binders and the like. See, e.g., Liberman et al., *Pharmaceutical Dosage Forms*, 2 Ed., Vol. 1, pp. 209-214 (1990); Singh et al., *Encyclopedia of Pharmaceutical Technology*, 2<sup>nd</sup> Ed., pp. 751-753 (2002); U.S. Pat. Nos. 4,327,725, 4,624,848, 4,968, 509, 5,461,140, 5,456,923, 5,516,527, 5,622,721, 5,686,105, 5,700,410, 5,977,175, 6,465,014 and 6,932,983, each of which is specifically incorporated by reference.

In some embodiments, pharmaceutical formulations are provided that include particles of Compound 1 and at least one dispersing agent or suspending agent for oral administration to a subject. The formulations may be a powder and/or granules for suspension, and upon admixture with water, a substantially uniform suspension is obtained.

It is to be appreciated that there is overlap between the above-listed additives used in the aqueous dispersions or suspensions described herein, since a given additive is often classified differently by different practitioners in the field, or is commonly used for any of several different functions. Thus, the above-listed additives should be taken as merely exemplary, and not limiting, of the types of additives that can be included in formulations described herein. The amounts of such additives can be readily determined by one skilled in the art, according to the particular properties desired.

Dosing and Treatment Regimens

In some embodiments, the amount of Compound 1 that is administered to a mammal is from 300 mg/day up to, and including, 1000 mg/day. In some embodiments, the amount of Compound 1 that is administered to a mammal is from 420 mg/day up to, and including, 840 mg/day. In some embodiments, the amount of Compound 1 that is administered to a mammal is about 420 mg/day, about 560 mg/day, or about 840 mg/day. In some embodiments, the amount of Compound 1 that is administered to a mammal is about 420 mg/day. In some embodiments, the amount of Compound 1 that is administered to a mammal is about 560 mg/day. In some embodiments, the AUC<sub>0-24</sub> of Compound 1 is between about 150 and about 3500 ng\*h/mL. In some embodiments, the AUC<sub>0-24</sub> of Compound 1 is between about 500 and about 1100 ng\*h/mL. In some embodiments, Compound 1 is administered orally. In some embodiments, Compound 1 is administered once per day, twice per day, or three times per day. In some embodiments, Compound 1 is administered daily. In some embodiments, Compound 1 is administered once daily. In some embodiments, Compound 1 is administered every other day. In some embodiments, the Compound 1 is a maintenance therapy.

Compound 1 can be used in the preparation of medicaments for the inhibition of Btk or a homolog thereof, or for the treatment of diseases or conditions that would benefit, at least in part, from inhibition of Btk or a homolog thereof, including a subject diagnosed with a hematological malignancy. In addition, a method for treating any of the diseases or conditions described herein in a subject in need of such treatment, involves administration of pharmaceutical compositions containing Compound 1, or a pharmaceutically acceptable salt, pharmaceutically acceptable N-oxide, pharmaceutically active metabolite, pharmaceutically acceptable prodrug, or pharmaceutically acceptable solvate thereof, in therapeutically effective amounts to said subject.

The compositions containing Compound 1 can be administered for prophylactic, therapeutic, or maintenance treatment. In some embodiments, compositions containing Compound 1 are administered for therapeutic applications (e.g., administered to a subject diagnosed with a hematological

malignancy). In some embodiments, compositions containing Compound 1 are administered for therapeutic applications (e.g., administered to a subject susceptible to or otherwise at risk of developing a hematological malignancy). In some embodiments, compositions containing Compound 1 are administered to a patient who is in remission as a maintenance therapy.

Amounts of Compound 1 will depend on the use (e.g., therapeutic, prophylactic, or maintenance). Amounts of Compound 1 will depend on severity and course of the disease or condition, previous therapy, the patient's health status, weight, and response to the drugs, and the judgment of the treating physician. It is considered well within the skill of the art for one to determine such therapeutically effective amounts by routine experimentation (including, but not limited to, a dose escalation clinical trial). In some embodiments, the amount of Compound 1 is from 300 mg/day up to, and including, 1000 mg/day. In some embodiments, the amount of Compound 1 is from 420 mg/day up 20 to, and including, 840 mg/day. In some embodiments, the amount of Compound 1 is from 400 mg/day up to, and including, 860 mg/day. In some embodiments, the amount of Compound 1 is about 360 mg/day. In some embodiments, the amount of Compound 1 is about 420 mg/day. In some 25 embodiments, the amount of Compound 1 is about 560 mg/day. In some embodiments, the amount of Compound 1 is about 840 mg/day. In some embodiments, the amount of Compound 1 is from 2 mg/kg/day up to, and including, 13 mg/kg/day. In some embodiments, the amount of Compound 1 is from 2.5 mg/kg/day up to, and including, 8 mg/kg/day. In some embodiments, the amount of Compound 1 is from 2.5 mg/kg/day up to, and including, 6 mg/kg/day. In some embodiments, the amount of Compound 1 is from 2.5 mg/kg/day up to, and including, 4 mg/kg/day. In some 35 embodiments, the amount of Compound 1 is about 2.5 mg/kg/day. In some embodiments, the amount of Compound 1 is about 8 mg/kg/day.

In one embodiment, the tablet formulation of the invention with 140 mg of dose in dogs produces  $C_{max}$  of about 260 40 to 400 ng/mL (Fed), and about 300 to 400 ng/mL (Fasted). In another embodiment, the formulation produces  $C_{max}$  of about 280 to 380 ng/mL (Fed), and about 360 to 380 ng/mL (Fasted). In a particular embodiment, the formulation produces  $C_{max}$  of about 290 ng/mL (Fed), and about 370 ng/mL (Fasted). In another particular embodiment, the formulation produces  $C_{max}$  of about 370 ng/mL (Fed), and about 370 ng/mL (Fasted). In one embodiment, the formulation is a wet granulation formulation. In one embodiment, the tablet formulation is Formulation BK02, BK21A, or BK21B. In a 50 particular embodiment, the tablet formulation is Formulation BK21A. In another particular embodiment, the tablet formulation is Formulation BK21B. (Table 1E and 1F).

In one embodiment, the tablet formulation of the invention with 140 mg of dose in dogs produces AUC of about 55 850 to 1050 ng\*h/mL (Fed), and about 850 to 1050 ng\*h/mL (Fasted). In another embodiment, the formulation produces AUC of about 870 to 1050 ng\*h/mL (Fed), and about 840 to 1000 ng\*h/mL (Fasted). In a particular embodiment, the formulation produces AUC of about 875 ng\*h/mL (Fed), 60 and about 1000 ng\*h/mL (Fasted). In another particular embodiment, the formulation produces AUC of about 1000 ng\*h/mL (Fed), and about 850 ng\*h/mL (Fasted). In one embodiment, the formulation is a wet granulation formulation. In one embodiment, the tablet formulation is Formulation BK02, BK21A, or BK21B. In a particular embodiment, the tablet formulation BK21A. In

72

another particular embodiment, the tablet formulation is Formulation BK21B. (Table 1E and 1F).

In one embodiment, the tablet formulation of the invention with 140 mg of dose in dogs produces %  $F_{rel}$  (tablet/capsule) ( $C_{max}$ ) value of about 150-250 (Fed) and 100-160 (Fasted). In a particular embodiment, the formulation produces %  $F_{rel}$  (tablet/capsule) ( $C_{max}$ ) value of about 170 (Fed) and about 110 (Fasted). In another particular embodiment, the formulation produces %  $F_{rel}$  (tablet/capsule) ( $C_{max}$ ) value of about 230 (Fed) and about 150 (Fasted). In one embodiment, the formulation is a wet granulation formulation. In one embodiment, the tablet formulation is Formulation BK02, BK21A, or BK21B. In a particular embodiment, the tablet formulation BK21A. In another particular embodiment, the tablet formulation is Formulation BK21B. (Table 1E and 1F).

In one embodiment, the tablet formulation of the invention with 140 mg of dose in dogs produces %  $F_{rel}$  (tablet/capsule) (AUC) value of about 110-150 (Fed) and 100-140 (Fasted). In a particular embodiment, the formulation produces %  $F_{rel}$  (tablet/capsule) (AUC) value of about 120 (Fed) and about 110 (Fasted). In another particular embodiment, the formulation produces %  $F_{rel}$  (tablet/capsule) (AUC) value of about 150 (Fed) and about 130 (Fasted). In one embodiment, the formulation is a wet granulation formulation. In one embodiment, the tablet formulation is Formulation BK02, BK21A, or BK21B. In a particular embodiment, the tablet formulation is Formulation BK21A. In another particular embodiment, the tablet formulation is Formulation BK21B. (Table 1E and 1F).

In one embodiment, the tablet formulation of the invention with 140 mg of dose in dogs produces %  $F_{rel}$  (Fed/Fasted) ( $C_{max}$ ) value of about 90-105. In a particular embodiment, the formulation produces %  $F_{rel}$  (Fed/Fasted) ( $C_{max}$ ) value of about 95. In another particular embodiment, the formulation produces %  $F_{rel}$  (Fed/Fasted) ( $C_{max}$ ) value of about 100. In one embodiment, the formulation is a wet granulation formulation. In one embodiment, the tablet formulation is Formulation BK02, BK21A, or BK21B. In a particular embodiment, the tablet formulation is Formulation BK21A. In another particular embodiment, the tablet formulation is Formulation BK21B. (Table 1E and 1F).

In one embodiment, the tablet formulation of the invention with 140 mg of dose in dogs produces %  $F_{rel}$  (Fed/Fasted) (AUC) value of about 90-140. In a particular embodiment, the formulation produces %  $F_{rel}$  (Fed/Fasted) (AUC) value of about 100. In one embodiment, the formulation is a wet granulation formulation. In one embodiment, the tablet formulation is Formulation BK21A, or BK21B. In a particular embodiment, the tablet formulation is Formulation BK21A. In another particular embodiment, the tablet formulation is Formulation BK21B. (Table 1E and 1F).

In some embodiments, pharmaceutical compositions described herein include about 140 mg of Compound 1. In some embodiments, a tablet formulation is prepared that includes about 140 mg of Compound 1. In some embodiments, 2, 3, 4, or 5 of the tablet formulations are administered daily. In some embodiments, 3 or 4 of the capsules are administered once daily. In some embodiments, the capsules are administered once daily. In other embodiments, the tablet are administered multiple times a day.

In another aspect is a high-load solid tablet formulation comprising ibrutinib, wherein ibrutinib is a compound with the structure of Compound 1, Compound 1

and the tablet comprises about 560 mg of ibrutinib.

In another embodiment is a high-load solid tablet formulation, wherein the tablet is used for once a day oral dosing. The high-load solid tablet formulations described herein make it possible for one tablet a day administration and contain a large amount of ibrutinib per tablet of about 420 mg to about 840 mg, such as about 420 mg, about 560 mg, or about 840 mg, or any range between any two of the values, end points inclusive. In another embodiment is a high-load solid tablet formulation, wherein the tablet comprises 560 mg of ibrutinib. In another embodiment is a high-load solid tablet formulation, wherein ibrutinib is in micronized form

In some embodiments, Compound 1 is administered daily. 35 In some embodiments, Compound 1 is administered every other day.

In some embodiments, Compound 1 is administered once per day. In some embodiments, Compound 1 is administered twice per day. In some embodiments, Compound 1 is 40 administered three times per day. In some embodiments, Compound 1 is administered four times per day.

In some embodiments, Compound 1 is administered until disease progression, unacceptable toxicity, or individual choice. In some embodiments, Compound 1 is administered 45 daily until disease progression, unacceptable toxicity, or individual choice. In some embodiments, Compound 1 is administered every other day until disease progression, unacceptable toxicity, or individual choice.

In the case wherein the patient's status does improve, 50 upon the doctor's discretion the administration of the compounds may be given continuously; alternatively, the dose of drug being administered may be temporarily reduced or temporarily suspended for a certain length of time (i.e., a "drug holiday"). The length of the drug holiday can vary 55 between 2 days and 1 year, including by way of example only, 2 days, 3 days, 4 days, 5 days, 6 days, 7 days, 10 days, 12 days, 15 days, 20 days, 28 days, 35 days, 50 days, 70 days, 100 days, 120 days, 150 days, 180 days, 200 days, 250 days, 280 days, 300 days, 320 days, 350 days, or 365 days. The dose reduction during a drug holiday may be from 10%-100%, including, by way of example only, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, or 100%.

Once improvement of the patient's conditions has 65 occurred, a maintenance dose is administered if necessary. Subsequently, the dosage or the frequency of administration,

or both, can be reduced, as a function of the symptoms, to a level at which the improved disease, disorder or condition is retained. Patients can, however, require intermittent treatment on a long-term basis upon any recurrence of symptoms.

The amount of a given agent that will correspond to such an amount will vary depending upon factors such as the particular compound, the severity of the disease, the identity (e.g., weight) of the subject or host in need of treatment, but can nevertheless be routinely determined in a manner known in the art according to the particular circumstances surrounding the case, including, e.g., the specific agent being administered, the route of administration, and the subject or host being treated. In general, however, doses employed for adult human treatment will typically be in the range of 0.02-5000 mg per day, or from about 1-1500 mg per day. The desired dose may conveniently be presented in a single dose or as divided doses administered simultaneously (or over a short period of time) or at appropriate intervals, for example as two, three, four or more sub-doses per day.

The pharmaceutical compositions or formulations described herein may be in unit dosage forms suitable for single administration of precise dosages. In unit dosage form, the formulation is divided into unit doses containing appropriate quantities of one or more compound. The unit dosage may be in the form of a package containing discrete quantities of the formulation. Non-limiting examples are packaged tablets or capsules, and powders in vials or ampoules. Aqueous suspension compositions can be packaged in single-dose non-reclosable containers. Alternatively, multiple-dose reclosable containers can be used, in which case it is typical to include a preservative in the composition. In some embodiments, each unit dosage form comprises 140 mg of Compound 1. In some embodiments, an individual is administered 1 unit dosage form per day. In some embodiments, an individual is administered 2 unit dosage forms per day. In some embodiments, an individual is administered 3 unit dosage forms per day. In some embodiments, an individual is administered 4 unit dosage forms per day.

The foregoing ranges are merely suggestive, as the number of variables in regard to an individual treatment regime is large, and considerable excursions from these recommended values are not uncommon. Such dosages may be altered depending on a number of variables, not limited to the activity of the compound used, the disease or condition to be treated, the mode of administration, the requirements of the individual subject, the severity of the disease or condition being treated, and the judgment of the practitioner.

Toxicity and therapeutic efficacy of such therapeutic regimens can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, including, but not limited to, the determination of the  $LD_{50}$  (the dose lethal to 50% of the population) and the  $ED_{50}$  (the dose therapeutically effective in 50% of the population). The dose ratio between the toxic and therapeutic effects is the therapeutic index and it can be expressed as the ratio between LD<sub>50</sub> and ED<sub>50</sub>. Compounds exhibiting high therapeutic indices are preferred. The data obtained from cell culture assays and animal studies can be used in formulating a range of dosage for use in human. The dosage of such compounds lies preferably within a range of circulating concentrations that include the ED<sub>50</sub> with minimal toxicity. The dosage may vary within this range depending upon the dosage form employed and the route of administration utilized. Combination Therapy

In certain instances, it is appropriate to administer Compound 1 in combination with another therapeutic agent.

In one embodiment, the compositions and methods described herein are also used in conjunction with other therapeutic reagents that are selected for their particular usefulness against the condition that is being treated. In general, the compositions described herein and, in embodiments where combinational therapy is employed, other agents do not have to be administered in the same pharmaceutical composition, and are, because of different physical and chemical characteristics, administered by different routes. In one embodiment, the initial administration is made according to established protocols, and then, based upon the observed effects, the dosage, modes of administration and times of administration, further modified.

In various embodiments, the compounds are administered 15 concurrently (e.g., simultaneously, essentially simultaneously or within the same treatment protocol) or sequentially, depending upon the nature of the disease, the condition of the patient, and the actual choice of compounds used. In certain embodiments, the determination of the order of 20 administration, and the number of repetitions of administration of each therapeutic agent during a treatment protocol, is based upon evaluation of the disease being treated and the condition of the patient.

For combination therapies described herein, dosages of 25 the co-administered compounds vary depending on the type of co-drug employed, on the specific drug employed, on the disease or condition being treated and so forth.

The individual compounds of such combinations are administered either sequentially or simultaneously in separate or combined pharmaceutical formulations. In one embodiment, the individual compounds will be administered simultaneously in a combined pharmaceutical formulation. Appropriate doses of known therapeutic agents will be 35 men comprises dexamethasone and lenalidomide. appreciated by those skilled in the art.

The combinations referred to herein are conveniently presented for use in the form of a pharmaceutical compositions together with a pharmaceutically acceptable diluent (s) or carrier(s).

Disclosed herein, in certain embodiments, is a method for treating a cancer in an individual in need thereof, comprising: administering to the individual an amount of Compound 1. In some embodiments, the method further comprises administering a second cancer treatment regimen.

In some embodiments, administering a Btk inhibitor before a second cancer treatment regimen reduces immunemediated reactions to the second cancer treatment regimen. In some embodiments, administering Compound 1 before ofatumumab reduces immune-mediated reactions to ofatu- 50 mumah.

In some embodiments, the second cancer treatment regimen comprises a chemotherapeutic agent, a steroid, an immunotherapeutic agent, a targeted therapy, or a combination thereof. In some embodiments, the second cancer 55 treatment regimen comprises a B cell receptor pathway inhibitor. In some embodiments, the B cell receptor pathway inhibitor is a CD79A inhibitor, a CD79B inhibitor, a CD19 inhibitor, a Lyn inhibitor, a Syk inhibitor, a PI3K inhibitor, a Blnk inhibitor, a PLCγ inhibitor, a PKCβ inhibitor, or a 60 combination thereof. In some embodiments, the second cancer treatment regimen comprises an antibody, B cell receptor signaling inhibitor, a PI3K inhibitor, an IAP inhibitor, an mTOR inhibitor, an immunochemotherapy, a radioimmunotherapeutic, a DNA damaging agent, a proteosome inhibitor, a Cyp3A4 inhibitor, a histone deacetylase inhibitor, a protein kinase inhibitor, a hedgehog inhibitor, an

76

Hsp90 inhibitor, a telomerase inhibitor, a Jak1/2 inhibitor, a protease inhibitor, a PKC inhibitor, a PARP inhibitor, or a combination thereof.

In some embodiments, the second cancer treatment regimen comprises chlorambucil, ifosphamide, doxorubicin, mesalazine, thalidomide, lenalidomide, temsirolimus, everolimus, fludarabine, fostamatinib, paclitaxel, docetaxel, ofatumumab, rituximab, dexamethasone, prednisone, CAL-101, ibritumomab, tositumomab, bortezomib, pentostatin, endostatin, EPOCH-R, DA-EPOCH-R, rifampin, selinexor, gemcitabine, obinutuzumab, carmustine, cytarabine, melphalan, ublituximab, palbociclib, ACP-196 (Acerta Pharma BV), TGR-1202 (TG Therapeutics, Inc.), TEDDI, TEDD, MEDI4736 (AstraZeneca), ABT-0199 (AbbVie), CC-122 (Celgene Corporation), LD-AraC, ketoconazole, etoposide, carboplatin, moxifloxacin, citrovorum, methotrexate, filgrastim, mesna, vincristine, cyclophosphamide, erythromycin, voriconazole, nivolumab, or a combination thereof.

In some embodiments, the second cancer treatment regimen comprises cyclophosphamide, hydroxydaunorubicin, vincristine, and prednisone, and optionally, rituximab.

In some embodiments, the second cancer treatment regimen comprises bendamustine, and rituximab.

In some embodiments, the second cancer treatment regimen comprises fludarabine, cyclophosphamide, and rituximab.

In some embodiments, the second cancer treatment regimen comprises cyclophosphamide, vincristine, and prednisone, and optionally, rituximab.

In some embodiments, the second cancer treatment regimen comprises etoposide, doxorubicin, vinristine, cyclophosphamide, prednisolone, and optionally, rituximab.

In some embodiments, the second cancer treatment regi-

In some embodiments, the second cancer treatment comprises a proteasome inhibitor. In some embodiments, the second treatment comprises bortezomib. In some embodiments, the second cancer treatment comprises an epoxyketone. In some embodiments, the second cancer treatment comprises epoxomicin. In some embodiments, the second cancer treatment comprises a tetrapeptide epoxyketone In some embodiments, the second cancer treatment comprises carfilzomib. In some embodiments, the second cancer treat-45 ment comprises disulfram, epigallocatechin-3-gallate, salinosporamide A, ONX 0912m CEP-18770, MLN9708, or MG132.

In some embodiments, the second cancer treatment comprises a Cyp3A4 inhibitor. In some embodiments, the second cancer treatment comprises indinavir, nelfinavir, ritonavir, clarithromycin, itraconazole, ketoconazole, nefazodone. In some embodiments, the second cancer treatment comprises ketoconazole.

In some embodiments, the second cancer treatment comprises a Janus Kinase (JAK) inhibitor. In some embodiments, the second treatment comprises Lestaurtinib, Tofacitinib, Ruxolitinib, CYT387, Baricitinib or Pacritinib.

In some embodiments, the second cancer treatment comprises a histone deacetylase inhibitor (HDAC inhibitor, HDI). In some embodiments, the second cancer treatment comprises a hydroxamic acid (or hydroxamate), such as trichostatin A, vorinostat (SAHA), belinostat (PXD101), LAQ824, and panobinostat (LBH589), a cyclic tetrapeptide, such as trapoxin B, a depsipeptide, a benzamide, such as entinostat (MS-275),CI994, and mocetinostat (MGCD0103), an electrophilic ketone, or an aliphatic acid compound, such as phenylbutyrate and valproic acid,

Additional cancer treatment regimens include Nitrogen Mustards such as for example, bendamustine, chlorambucil, chlormethine, cyclophosphamide, ifosfamide, melphalan, prednimustine, trofosfamide; Alkyl Sulfonates like busulfan, mannosulfan, treosulfan; Ethylene Imines like carboquone, 5 thiotepa, triaziquone; Nitrosoureas like carmustine, fotemustine, lomustine, nimustine, ranimustine, semustine, streptozocin; Epoxides such as for example, etoglucid; Other Alkylating Agents such as for example dacarbazine, mitobronitol, pipobroman, temozolomide; Folic Acid Ana- 10 logues such as for example methotrexate, permetrexed, pralatrexate, raltitrexed; Purine Analogs such as for example cladribine, clofarabine, fludarabine, mercaptopurine, nelarabine, tioguanine; Pyrimidine Analogs such as for example azacitidine, capecitabine, carmofur, cytarabine, decitabine, 15 fluorouracil, gemcitabine, tegafur; Vinca Alkaloids such as for example vinblastine, vincristine, vindesine, vinflunine, vinorelbine; Podophyllotoxin Derivatives such as for example etoposide, teniposide; Colchicine derivatives such as for example demecolcine; Taxanes such as for example 20 docetaxel, paclitaxel, paclitaxel poliglumex; Other Plant Alkaloids and Natural Products such as for example trabectedin; Actinomycines such as for example dactinomycin; Antracyclines such as for example aclarubicin, daunorubicin, doxorubicin, epirubicin, idarubicin, mitoxantrone, pira- 25 rubicin, valrubicin, zorubincin; Other Cytotoxic Antibiotics such as for example bleomycin, ixabepilone, mitomycin, plicamycin; Platinum Compounds such as for example carboplatin, cisplatin, oxaliplatin, satraplatin; Methylhydrazines such as for example procarbazine; Sensitizers such as 30 for example aminolevulinic acid, efaproxiral, methyl aminolevulinate, porfimer sodium, temoporfin; Protein Kinase Inhibitors such as for example dasatinib, erlotinib, everolimus, gefitinib, imatinib, lapatinib, nilotinib, pazonanib, sorafenib, sunitinib, temsirolimus; Other Antineoplastic 35 Agents such as for example alitretinoin, altretamine, amzacrine, anagrelide, arsenic trioxide, asparaginase, bexarotene, bortezomib, celecoxib, denileukin diftitox, estramustine, hydroxycarbamide, irinotecan, lonidamine, masoprocol, miltefosein, mitoguazone, mitotane, oblimersen, pegaspar- 40 gase, pentostatin, romidepsin, sitimagene ceradenovec, tiazofurine, topotecan, tretinoin, vorinostat; Estrogens such as for example diethylstilbenol, ethinylestradiol, fosfestrol, polyestradiol phosphate; Progestogens such as for example gestonorone, medroxyprogesterone, megestrol; Gonadotro- 45 pin Releasing Hormone Analogs such as for example buserelin, goserelin, leuprorelin, triptorelin; Anti-Estrogens such as for example fulvestrant, tamoxifen, toremifene; Anti-Androgens such as for example bicalutamide, flutamide, nilutamide; Enzyme Inhibitors, aminoglutethimide, anastro- 50 zole, exemestane, formestane, letrozole, vorozole; Other Hormone Antagonists such as for example abarelix, degarelix; Immunostimulants such as for example histamine dihydrochloride, mifamurtide, pidotimod, plerixafor, roquinimex, thymopentin; Immunosuppressants such as for 55 example everolimus, gusperimus, leflunomide, mycophenolic acid, sirolimus; Calcineurin Inhibitors such as for example ciclosporin, tacrolimus; Other Immunosuppressants such as for example azathioprine, lenalidomide, methotrexate, thalidomide; and Radiopharmaceuticals such 60

Additional cancer treatment regimens include interferons, interleukins, Tumor Necrosis Factors, Growth Factors, or the like.

as for example, iobenguane.

Additional cancer treatment regimens include Immunos- 65 timulants such as for example ancestim, filgrastim, lenograstim, molgramostim, pegfilgrastim, sargramostim;

78

Interferons such as for example interferon alfa natural, interferon alfa-2a, interferon alfa-2b, interferon alfacon-1, interferon alfa-n1, interferon beta natural, interferon beta-1a, interferon beta-1b, interferon gamma, peginterferon alfa-2a, peginterferon alfa-2b; Interleukins such as for example aldesleukin, oprelvekin; Other Immunostimulants such as for example BCG vaccine, glatiramer acetate, histamine dihydrochloride, immunocyanin, lentinan, melanoma vaccine, mifamurtide, pegademase, pidotimod, plerixafor, poly I:C, poly ICLC, roquinimex, tasonermin, thymopentin; Immunosuppressants such as for example abatacept, abetimus, alefacept, antilymphocyte immunoglobulin (horse), antithymocyte immunoglobulin (rabbit), eculizumab, efalizumab, everolimus, gusperimus, leflunomide, muromab-CD3. mycophenolic acid, natalizumab, sirolimus; TNF alpha Inhibitors such as for example adalimumab, afelimomab, certolizumab pegol, etanercept, golimumab, infliximab; Interleukin Inhibitors such as for example anakinra, basiliximab, canakinumab, daclizumab, mepolizumab, rilonacept, tocilizumab, ustekinumab; Calcineurin Inhibitors such as for example ciclosporin, tacrolimus; Other Immunosuppressants such as for example azathioprine, lenalidomide, methotrexate, thalidomide.

Additional cancer treatment regimens include Adalimumab, Alemtuzumab, Basiliximab, Bevacizumab, Cetuximab, Certolizumab pegol, Daclizumab, Eculizumab, Efalizumab, Gemtuzumab, Ibritumomab tiuxetan, Infliximab, Muromonab-CD3, Natalizumab, Panitumumab, Ranibizumab, Rituximab, Tositumomab, Trastuzumab, or the like, or a combination thereof.

Additional cancer treatment regimens include Monoclonal Antibodies such as for example alemtuzumab, bevacizumab, catumaxomab, cetuximab, edrecolomab, gemtupanitumumab, ofatumumab, trastuzumab, Immunosuppressants, eculizumab, efalizumab, muromab-CD3, natalizumab; TNF alpha Inhibitors such as for example adalimumab, afelimomab, certolizumab pegol, golimumab, infliximab, Interleukin Inhibitors, basiliximab, canakinumab, daclizumab, mepolizumab, tocilizumab, ustekinumab, Radiopharmaceuticals, ibritumomab tiuxetan, tositumomab; Others Monoclonal Antibodies such as for example abagovomab, adecatumumab, alemtuzumab, anti-CD30 monoclonal antibody Xmab2513, anti-MET monoclonal antibody MetMab, apolizumab, apomab, arcitubasiliximab, bispecific antibody momab. blinatumomab, brentuximab vedotin, capromab pendetide, cixutumumab, claudiximab, conatumumab, dacetuzumab, denosumab, eculizumab, epratuzumab, epratuzumab, ertumaxomab, etaracizumab, figitumumab, fresolimumab, galiximab, ganitumab, gemtuzumab ozogamicin, glembatumumab, ibritumomab, inotuzumab ozogamicin, ipilimumab, lexatumumab, lintuzumab, lucatumumab, mapatumumab, matuzumab, milatuzumab, monoclonal antibody CC49, necitumumab, nimotuzumab, ofatumumab, oregovomab, pertuzumab, ramacurimab, ranibizumab, siplizumab, sonepcizumab, tanezumab, tositumomab, trastuzumab, tremelimumab, tucotuzumab celmoleukin, veltuzumab, visilizumab, volociximab, zalutumumab.

Additional cancer treatment regimens include agents that affect the tumor micro-environment such as cellular signaling network (e.g. phosphatidylinositol 3-kinase (PI3K) signaling pathway, signaling from the B-cell receptor and the IgE receptor). In some embodiments, the second agent is a PI3K signaling inhibitor or a syc kinase inhibitor. In one embodiment, the syk inhibitor is R788. In another embodiment is a PKCy inhibitor such as by way of example only, enzastaurin.

In some embodiments, the additional therapeutic agent comprises an analgesic such as acetaminophen.

In some embodiments, the additional therapeutic agent comprises an agent selected from: an inhibitor of LYN, SYK, JAK, PI3K, PLCγ, MAPK, MEK or NFκB.

Examples of agents that affect the tumor micro-environment include PI3K signaling inhibitor, syc kinase inhibitor, Protein Kinase Inhibitors such as for example dasatinib, erlotinib, everolimus, gefitinib, imatinib, lapatinib, nilotinib, pazonanib, sorafenib, sunitinib, temsirolimus; Other Angio- 10 genesis Inhibitors such as for example GT-111, JI-101, R1530; Other Kinase Inhibitors such as for example AC220, AC480, ACE-041, AMG 900, AP24534, Arry-614, AT7519, AT9283, AV-951, axitinib, AZD1152, AZD7762, AZD8055, AZD8931, bafetinib, BAY 73-4506, BGJ398, BGT226, 15 BI811283, BI6727, BIBF 1120, BIBW 2992, BMS-690154, BMS-777607, BMS-863233, BSK-461364, CAL-101, CEP-11981, CYC116, DCC-2036, dinaciclib, dovitinib lactate, E7050, EMD 1214063, ENMD-2076, fostamatinib disodium, GSK2256098, GSK690693, INCB18424, INNO-406, 20 JNJ-26483327, JX-594, KX2-391, linifanib, LY2603618, MGCD265, MK-0457, MK1496, MLN8054, MLN8237, MP470, NMS-1116354, NMS-1286937, ON 01919.Na, OSI-027, OSI-930, Btk inhibitor, PF-00562271. PF-02341066, PF-03814735, PF-04217903, PF-04554878, 25 PF-04691502, PF-3758309, PHA-739358, PLC3397, progenipoietin, R547, R763, ramucirumab, regorafenib, RO5185426, SAR103168, SCH 727965, SGI-1176, SGX523, SNS-314, TAK-593, TAK-901, TKI258, TLN-232, TTP607, XL147, XL228, XL281RO5126766, XL418, 30 XL765.

Further examples of anti-cancer agents for use in combination with a Btk inhibitor compound include inhibitors of mitogen-activated protein kinase signaling, e.g., U0126, PD98059, PD184352, PD0325901, ARRY-142886, 35 SB239063, SP600125, BAY 43-9006, wortmannin, or LY294002; Syk inhibitors; mTOR inhibitors; and antibodies (e.g., rituxan).

Other anti-cancer agents that can be employed in combination with a Btk inhibitor compound include Adriamycin, 40 Dactinomycin, Bleomycin, Vinblastine, Cisplatin, acivicin; aclarubicin; acodazole hydrochloride; acronine; adozelesin; aldesleukin; altretamine; ambomycin; ametantrone acetate; aminoglutethimide; amsacrine; anastrozole; anthramycin; asparaginase; asperlin; azacitidine; azetepa; azotomycin; 45 batimastat; benzodepa; bicalutamide; bisantrene hydrochloride; bisnafide dimesvlate; bizelesin; bleomycin sulfate; brequinar sodium; bropirimine; busulfan; cactinomycin; calusterone; caracemide; carbetimer; carboplatin; carmustine; carubicin hydrochloride; carzelesin; cedefingol; 50 chlorambucil; cirolemycin; cladribine; crisnatol mesylate; cyclophosphamide; cytarabine; dacarbazine; daunorubicin hydrochloride; decitabine; dexormaplatin; dezaguanine; dezaguanine mesylate; diaziquone; doxorubicin; doxorubicin hydrochloride; droloxifene; droloxifene citrate; dromo- 55 stanolone propionate; duazomycin; edatrexate; eflomithine hydrochloride; elsamitrucin; enloplatin; enpromate; epipropidine; epirubicin hydrochloride; erbulozole; esorubicin hydrochloride; estramustine; estramustine phosphate sodium; etanidazole; etoposide; etoposide phosphate; eto- 60 prine; fadrozole hydrochloride; fazarabine; fenretinide; floxuridine; fludarabine phosphate; fluorouracil; flurocitabine; fosquidone; fostriecin sodium; gemcitabine; gemcitabine hydrochloride; hydroxyurea; idarubicin hydrochloride; ifosfamide; iimofosine; interleukin Il (including 65 recombinant interleukin II, or r1L2), interferon alfa-2a; interferon alfa-2b; interferon alfa-n1; interferon alfa-n3; inter80

feron beta-1 a; interferon gamma-1 b; iproplatin; irinotecan hydrochloride; lanreotide acetate; letrozole; leuprolide acetate; liarozole hydrochloride; lometrexol sodium; lomustine; losoxantrone hydrochloride; masoprocol; maytansine; mechlorethamine hydrochloride; megestrol acetate; melengestrol acetate; melphalan; menogaril; mercaptopurine; methotrexate; methotrexate sodium; metoprine; meturedepa; mitindomide; mitocarcin; mitocromin; mitogillin; mitomalcin; mitomycin; mitosper; mitotane; mitoxantrone hydrochloride; mycophenolic acid; nocodazoie; nogalamycin; ormaplatin; oxisuran; pegaspargase; peliomycin; pentamustine; peplomycin sulfate; perfosfamide; pipobroman; piposulfan; piroxantrone hydrochloride; plicamyplomestane; porfimer sodium; porfiromycin; prednimustine; procarbazine hydrochloride; puromycin; puromycin hydrochloride; pyrazofurin; riboprine; rogletimide; safingol; safingol hydrochloride; semustine; simtrazene; sparfosate sodium; sparsomycin; spirogermanium hydrochloride; spiromustine; spiroplatin; streptonigrin; streptozocin; sulofenur; talisomycin; tecogalan sodium; tegafur; teloxantrone hydrochloride; temoporfin; teniposide; teroxirone; testolactone; thiamiprine; thioguanine; thiotepa; tiazofurin; tirapazamine; toremifene citrate; trestolone acetate; triciribine phosphate; trimetrexate; trimetrexate glucuronate; triptorelin; tubulozole hydrochloride; uracil mustard; uredepa; vapreotide; verteporfin; vinblastine sulfate; vincristine sulfate; vindesine; vindesine sulfate; vinepidine sulfate; vinglycinate sulfate; vinleurosine sulfate; vinorelbine tartrate; vinrosidine sulfate; vinzolidine sulfate; vorozole; zeniplatin; zinostatin; zorubicin hydrochloride.

Other anti-cancer agents that can be employed in combination with a Btk inhibitor compound include: 20-epi-1, 25 dihydroxyvitamin D3; 5-ethynyluracil; abiraterone; aclarubicin; acylfulvene; adecypenol; adozelesin; aldesleukin; ALL-TK antagonists; altretamine; ambamustine; amidox; amifostine; aminolevulinic acid; amrubicin; amsacrine; anagrelide; anastrozole; andrographolide; angiogenesis inhibitors; antagonist D; antagonist G; antarelix; anti-dorsalizing morphogenetic protein-1; antiandrogen, prostatic carcinoma; antiestrogen; antineoplaston; antisense oligonucleotides; aphidicolin glycinate; apoptosis gene modulators; apoptosis regulators; apurinic acid; ara-CDP-DL-PTBA; arginine deaminase; asulacrine; atamestane; atrimustine; axinastatin 1; axinastatin 2; axinastatin 3; azasetron; azatoxin; azatyrosine; baccatin III derivatives; balanol; batimastat; BCR/ABL antagonists; benzochlorins; benzoylstaurosporine; beta lactam derivatives; beta-alethine; betaclamycin B; betulinic acid; bFGF inhibitor; bicalutamide; bisantrene; bisaziridinylspermine; bisnafide; bistratene A; bizelesin; breflate; bropirimine; budotitane; buthionine sulfoximine; calcipotriol; calphostin C; camptothecin derivatives; canarypox IL-2; capecitabine; carboxamide-aminotriazole; carboxyamidotriazole; CaRest M3; CARN 700; cartilage derived inhibitor; carzelesin; casein kinase inhibitors (ICOS); castanospermine; cecropin B; cetrorelix; chlorlns; chloroquinoxaline sulfonamide; cicaprost; cis-porphyrin; cladribine; clomifene analogues; clotrimazole; collismycin A; collismycin B; combretastatin A4; combretastatin analogue; conagenin; crambescidin 816; crisnatol; cryptophycin 8; cryptophycin A derivatives; curacin A; cyclopentanthraquinones; cycloplatam; cypemycin; cytarabine ocfosfate; cytolytic factor; cytostatin; dacliximab; decitabine; dehydrodidemnin B; deslorelin; dexamethasone; dexifosfamide; dexrazoxane; dexverapamil; diaziquone; didemnin B; didox; diethylnorspermine; dihydro-5-azacytidine; 9-dioxamycin; diphenyl spiromustine; docosanol; dolasetron; doxifluridine; droloxifene; dronabinol; duocar-

mycin SA; ebselen; ecomustine; edelfosine; edrecolomab; eflomithine; elemene; emitefur; epirubicin; epristeride; estramustine analogue; estrogen agonists; estrogen antagonists; etanidazole; etoposide phosphate; exemestane; fadrozole; fazarabine; fenretinide; filgrastim; finasteride; fla- 5 vopiridol; flezelastine; fluasterone; fludarabine; fluorodaunorunicin hydrochloride; forfenimex; formestane; fostriecin; fotemustine; gadolinium texaphyrin; gallium nitrate; galocitabine; ganirelix; gelatinase inhibitors; gemcitabine; glutathione inhibitors; hepsulfam; heregulin; hex- 10 amethylene bisacetamide; hypericin; ibandronic acid; idarubicin; idoxifene; idramantone; ilmofosine; ilomastat; imidazoacridones; imiquimod; immunostimulant peptides; insulin-such as for example growth factor-1 receptor inhibitor; interferon agonists; interferons; interleukins; ioben- 15 guane; iododoxorubicin; ipomeanol, 4-; iroplact; irsogladine; isobengazole; isohomohalicondrin B; itasetron; jasplakinolide; kahalalide F; lamellarin-N triacetate; lanreotide; leinamycin; lenograstim; lentinan sulfate; leptolstatin; letrozole; leukemia inhibiting factor; leukocyte alpha 20 interferon; leuprolide+estrogen+progesterone; leuprorelin; levamisole; liarozole; linear polyamine analogue; lipophilic disaccharide peptide; lipophilic platinum compounds; lissoclinamide 7; lobaplatin; lombricine; lometrexol; lonidamine; losoxantrone; lovastatin; loxoribine; lurtotecan; lutetium 25 texaphyrin; lysofylline; lytic peptides; maitansine; mannostatin A; marimastat; masoprocol; maspin; matrilysin inhibitors; matrix metalloproteinase inhibitors; menogaril; merbarone; meterelin; methioninase; metoclopramide; MIF inhibitor; mifepristone; miltefosine; mirimostim; mis- 30 matched double stranded RNA; mitoguazone; mitolactol; mitomycin analogues; mitonafide; mitotoxin fibroblast growth factor-saporin; mitoxantrone; mofarotene; molgramostim; monoclonal antibody, human chorionic gonadotrophin; monophosphoryl lipid A+myobacterium cell wall sk; 35 mopidamol; multiple drug resistance gene inhibitor; multiple tumor suppressor 1-based therapy; mustard anticancer agent; mycaperoxide B; mycobacterial cell wall extract; myriaporone; N-acetyldinaline; N-substituted benzamides; terpin; nartograstim; nedaplatin; nemorubicin; neridronic acid; neutral endopeptidase; nilutamide; nisamycin; nitric oxide modulators; nitroxide antioxidant; nitrullyn; O6-benzylguanine; octreotide; okicenone; oligonucleotides; onapristone; ondansetron; oracin; oral 45 cytokine inducer; ormaplatin; osaterone; oxaliplatin; oxaunomycin; palauamine; palmitoylrhizoxin; pamidronic acid; panaxytriol; panomifene; parabactin; pazelliptine; pegaspargase; peldesine; pentosan polysulfate sodium; pentostatin; pentrozole; perflubron; perfosfamide; perillyl alco- 50 hol; phenazinomycin; phenylacetate; phosphatase inhibitors; picibanil; pilocarpine hydrochloride; pirarubicin; piritrexim; placetin A; placetin B; plasminogen activator inhibitor; platinum complex; platinum compounds; platinum-triamine complex; porfimer sodium; porfiromycin; 55 prednisone; propyl bis-acridone; prostaglandin J2; proteasome inhibitors; protein A-based immune modulator; protein kinase C inhibitor; protein kinase C inhibitors, microalgal; protein tyrosine phosphatase inhibitors; purine nucleoside phosphorylase inhibitors; purpurins; pyrazolo- 60 acridine; pyridoxylated hemoglobin polyoxyethylerie conjugate; raf antagonists; raltitrexed; ramosetron; ras famesyl protein transferase inhibitors; ras inhibitors; ras-GAP inhibitor; retelliptine demethylated; rhenium Re 186 etidronate; rhizoxin; ribozymes; RII retinamide; rogletimide; rohi- 65 tukine; romurtide; roquinimex; rubiginone B1; ruboxyl; safingol; saintopin; SarCNU; sarcophytol A; sargramostim;

82

Sdi 1 mimetics; semustine; senescence derived inhibitor 1; sense oligonucleotides; signal transduction inhibitors; signal transduction modulators; single chain antigen-binding protein; sizofiran; sobuzoxane; sodium borocaptate; sodium phenylacetate; solverol; somatomedin binding protein; sonermin; sparfosic acid; spicamycin D; spiromustine; splenopentin; spongistatin 1; squalamine; stem cell inhibitor; stemcell division inhibitors; stipiamide; stromelysin inhibitors; sulfinosine; superactive vasoactive intestinal peptide antagonist; suradista; suramin; swainsonine; synthetic glycosaminoglycans; tallimustine; tamoxifen methiodide; tauromustine; tazarotene; tecogalan sodium; tegafur; tellurapyrylium; telomerase inhibitors; temoporfin; temozolomide; teniposide; tetrachlorodecaoxide; tetrazomine; thaliblastine; thiocoraline; thrombopoietin; thrombopoietin mimetic; thymalfasin; thymopoietin receptor agonist; thymotrinan; thyroid stimulating hormone; tin ethyl etiopurpurin; tirapazamine; titanocene bichloride; topsentin; toremifene; totipotent stem cell factor; translation inhibitors; tretinoin; triacetyluridine; triciribine; trimetrexate; triptorelin; tropisetron; turosteride; tyrosine kinase inhibitors; tyrphostins; UBC inhibitors; ubenimex; urogenital sinus-derived growth inhibitory factor; urokinase receptor antagonists; vapreotide; variolin B; vector system, erythrocyte gene therapy; velaresol; veramine; verdins; verteporfin; vinorelbine; vinxaltine; vitaxin; vorozole; zanoterone; zeniplatin; zilascorb; and zinostatin stimalamer.

Yet other anticancer agents that can be employed in combination with a Btk inhibitor compound include alkylating agents, antimetabolites, natural products, or hormones, e.g., nitrogen mustards (e.g., mechloroethamine, cyclophosphamide, chlorambucil, etc.), alkyl sulfonates (e.g., busulfan), nitrosoureas (e.g., carmustine, lomusitne, ete.), or triazenes (decarbazine, etc.). Examples of antimetabolites include but are not limited to folic acid analog (e.g., methotrexate), or pyrimidine analogs (e.g., Cytarabine), purine analogs (e.g., mercaptopurine, thioguanine, pentosta-

Examples of alkylating agents that can be employed in nafarelin; nagrestip; naloxone+pentazocine; napavin; naph- 40 combination a Btk inhibitor compound include, but are not limited to, nitrogen mustards (e.g., mechloroethamine, cyclophosphamide, chlorambucil, meiphalan, etc.), ethylenimine and methylmelamines (e.g., hexamethlymelamine, thiotepa), alkyl sulfonates (e.g., busulfan), nitrosoureas (e.g., carmustine, lomusitne, semustine, streptozocin, etc.), or triazenes (decarbazine, etc.). Examples of antimetabolites include, but are not limited to folic acid analog (e.g., methotrexate), or pyrimidine analogs (e.g., fluorouracil, floxouridine, Cytarabine), purine analogs (e.g., mercaptopurine, thioguanine, pentostatin.

Examples of hormones and antagonists include, but are not limited to, adrenocorticosteroids (e.g., prednisone), progestins (e.g., hydroxyprogesterone caproate, megestrol acetate, medroxyprogesterone acetate), estrogens (e.g., diethlystilbestrol, ethinyl estradiol), antiestrogen (e.g., tamoxifen), androgens (e.g., testosterone propionate, fluoxymesterone), antiandrogen (e.g., flutamide), gonadotropin releasing hormone analog (e.g., leuprolide).

Other agents that can be used in the methods and compositions described herein for the treatment or prevention of cancer include platinum coordination complexes (e.g., cisplatin, carboblatin), anthracenedione (e.g., mitoxantrone), substituted urea (e.g., hydroxyurea), methyl hydrazine derivative (e.g., procarbazine), adrenocortical suppressant (e.g., mitotane, aminoglutethimide).

Examples of anti-cancer agents which act by arresting cells in the G2-M phases due to stabilized microtubules and

**83** which can be used in combination with a Btk inhibitor

compound include without limitation the following marketed drugs and drugs in development: Erbulozole (also known as R-55104), Dolastatin 10 (also known as DLS-10 and NSC-376128), Mivobulin isethionate (also known as 5 CI-980), Vincristine, NSC-639829, Discodermolide (also known as NVP-XX-A-296), ABT-751 (Abbott, also known as E-7010), Altorhyrtins (such as Altorhyrtin A and Altorhyrtin C), Spongistatins (such as Spongistatin 1, Spongistatin 2, Spongistatin 3, Spongistatin 4, Spongistatin 5, 10 Spongistatin 6, Spongistatin 7, Spongistatin 8, and Spongistatin 9), Cemadotin hydrochloride (also known as LU-103793 and NSC-D-669356), Epothilones (such as Epothilone A, Epothilone B, Epothilone C (also known as desoxyepothilone A or dEpoA), Epothilone D (also referred 15 to as KOS-862, dEpoB, and desoxyepothilone B), Epothilone E, Epothilone F, Epothilone B N-oxide, Epothilone A N-oxide, 16-aza-epothilone B, 21-aminoepothilone B (also known as BMS-310705), 21-hydroxyepothilone D (also known as Desoxyepothilone F and dEpoF), 26-fluoroepoth- 20 ilone), Auristatin PE (also known as NSC-654663), Soblidotin (also known as TZT-1027), LS-4559-P (Pharmacia, also known as LS-4577), LS-4578 (Pharmacia, also known as LS-477-P), LS-4477 (Pharmacia), LS-4559 (Pharmacia), RPR-112378 (Aventis), Vincristine sulfate, DZ-3358 (Dai- 25 ichi), FR-182877 (Fujisawa, also known as WS-9885B), GS-164 (Takeda), GS-198 (Takeda), KAR-2 (Hungarian Academy of Sciences), BSF-223651 (BASF, also known as ILX-651 and LU-223651), SAH-49960 (Lilly/Novartis), SDZ-268970 (Lilly/Novartis), AM-97 (Armad/Kyowa 30 Hakko), AM-132 (Armad), AM-138 (Armad/Kyowa Hakko), IDN-5005 (Indena), Cryptophycin 52 (also known as LY-355703), AC-7739 (Ajinomoto, also known as AVE-8063A and CS-39.HCl), AC-7700 (Ajinomoto, also known as AVE-8062, AVE-8062A, CS-39-L-Ser.HCl, and RPR- 35 258062A), Vitilevuamide, Tubulysin A, Canadensol, Centaureidin (also known as NSC-106969), T-138067 (Tularik, also known as T-67, TL-138067 and TI-138067), COBRA-1 (Parker Hughes Institute, also known as DDE-261 and WHI-261), H10 (Kansas State University), H16 (Kansas 40 State University), Oncocidin A1 (also known as BTO-956 and DIME), DDE-313 (Parker Hughes Institute), Fijianolide B, Laulimalide, SPA-2 (Parker Hughes Institute), SPA-1 (Parker Hughes Institute, also known as SPIKET-P), 3-IAABU (Cytoskeleton/Mt. Sinai School of Medicine, also 45 known as MF-569), Narcosine (also known as NSC-5366), Nascapine, D-24851 (Asta Medica), A-105972 (Abbott), Hemiasterlin, 3-BAABU (Cytoskeleton/Mt. Sinai School of Medicine, also known as MF-191), TMPN (Arizona State University), Vanadocene acetylacetonate, T-138026 (Tu-50 larik), Monsatrol, Inanocine (also known as NSC-698666), 3-1AABE (Cytoskeleton/Mt. Sinai School of Medicine), A-204197 (Abbott), T-607 (Tularik, also known as T-900607), RPR-115781 (Aventis), Eleutherobins (such as Desmethyleleutherobin, Desaetyleleutherobin, Isoeleuther- 55 obin A, and Z-Eleutherobin), Caribaeoside, Caribaeolin, Halichondrin B, D-64131 (Asta Medica), D-68144 (Asta Medica), Diazonamide A, A-293620 (Abbott), NPI-2350 (Nereus), Taccalonolide A, TUB-245 (Aventis), A-259754 (Abbott), Diozostatin, (-)-Phenylahistin (also known as 60 NSCL-96F037), D-68838 (Asta Medica), D-68836 (Asta Medica), Myoseverin B, D-43411 (Zentaris, also known as D-81862), A-289099 (Abbott), A-318315 (Abbott), HTI-286 (also known as SPA-110, trifluoroacetate salt) (Wyeth), D-82317 (Zentaris), D-82318 (Zentaris), SC-12983 (NCI), 65 Resverastatin phosphate sodium, BPR-OY-007 (National Health Research Institutes), and SSR-250411 (Sanofi).

84

The formulations may be used in any combination with one or more other anti-thromboembolic agents to treat or prevent thromboembolic disorder (e.g., stroke). Examples of anti-thromboembolic agents include, but are not limited any of the following: thrombolytic agents (e.g., alteplase anistreplase, streptokinase, urokinase, or tissue plasminogen activator), heparin, tinzaparin, warfarin, dabigatran (e.g., dabigatran etexilate), factor Xa inhibitors (e.g., fondaparinux, draparinux, rivaroxaban, DX-9065a, otamixaban, LY517717, or YM150), ticlopidine, clopidogrel, CS-747 (prasugrel, LY640315), ximelagatran, or BIBR 1048.

In some embodiments, the additional anti-cancer agent that is a Bcl-2 inhibitor.

In some embodiments, the additional anti-cancer agent is immune checkpoint inhibitor. In some embodiments, the immune checkpoint inhibitor is an inhibitor of Programmed Death-Ligand 1 (PD-L1, also known as B7-H1, CD274), Programmed Death 1 (PD-1), CTLA-4, PD-L2 (B7-DC, CD273), LAG3, TIM3, 2B4, A2aR, B7H1, B7H3, B7H4, BTLA, CD2, CD27, CD28, CD30, CD40, CD70, CD80, CD86, CD137, CD160, CD226, CD276, DR3, GAL9, GITR, HAVCR2, HVEM, IDO1, IDO2, ICOS (inducible T cell costimulator), KIR, LAIR1, LIGHT, MARCO (macrophage receptor with collageneous structure), PS (phosphatidylserine), OX-40, SLAM, TIGHT, VISTA, VTCN1, or any combinations thereof. In some embodiments, the immune checkpoint inhibitor is an inhibitor of PD-L1, PD-1, CTLA-4, LAG3, or TIM3. In some embodiments, the immune checkpoint inhibitor is an inhibitor of PD-L1. In some embodiments, the immune checkpoint inhibitor is an inhibitor of PD-1. In some embodiments, the immune checkpoint inhibitor is an inhibitor of CTLA-4. In some embodiments, the immune checkpoint inhibitor is an inhibitor of LAG3. In some embodiments, the immune checkpoint inhibitor is an inhibitor of TIM3. In some embodiments, the immune checkpoint inhibitor is an inhibitor of PD-L2.

In some embodiments, the formulations are administered in combination with a CD20 inhibitor. Exemplary CD20 inhibitors include, but are not limited to, ibritumomab tiuxetan, ofatumumab, rituximab, tositumomab, and obinutuzumab

In some embodiments, the additional anticancer agent used in combination with the formulations described herein include CDK4 inhibitors (e.g., palbociclib).

In some embodiments, the additional cancer agent is a proteosome inhibitor. In some embodiments, the proteasome inhibitor is selected from bortezomib or carfilzomib

In some embodiments, the additional cancer agent that can be administered in combination with the formulations is an HDAC inhibitor. In some embodiments, the HDAC inhibitor is abexinostat or a salt thereof. In some embodiments, the abexinostat or a salt thereof is abexinostat HCl. In some embodiments, the abexinostat or a salt thereof is abexinostat tosylate.

In some embodiments, the additional cancer agent that can be administered in combination with the formulations is a MALT1 inhibitor, MCL-1 inhibitor, IDH1inhibitor, TLR inhibitor, or PIM inhibitor.

In some embodiments, the additional anti-cancer agent that can be administered in combination with the formulations is an immunomodulatory agent. Exemplary immunomodulatory agents include, but are not limited to, lenalidomide, thalidomide, and pomalidomide.

Where the individual is suffering from or at risk of suffering from an autoimmune disease, an inflammatory disease, or an allergy disease, Compound 1 can be used in with one or more of the following therapeutic agents in any

immunosuppressants (e.g., combination: tacrolimus, cyclosporin, rapamycin, methotrexate, cyclophosphamide, azathioprine, mercaptopurine, mycophenolate, or FTY720), glucocorticoids (e.g., prednisone, cortisone acetate, prednisolone, methylprednisolone, dexamethasone, betametha- <sup>5</sup> sone, triamcinolone, beclometasone, fludrocortisone acetate, deoxycorticosterone acetate, aldosterone), non-steroidal anti-inflammatory drugs (e.g., salicylates, arylalkanoic acids, 2-arylpropionic acids, N-arylanthranilic acids, oxicams, coxibs, or sulphonanilides), Cox-2-specific inhibitors (e.g., valdecoxib, celecoxib, or rofecoxib), leflunomide, gold thioglucose, gold thiomalate, aurofin, sulfasalazine, hydroxychloroquinine, minocycline, TNF-a binding proteins (e.g., infliximab, etanercept, or adalimumab), abatacept, anakinra, interferon-β, interferon-γ, interleukin-2, allergy vaccines, antihistamines, antileukotrienes, beta-agonists, theophylline, or anticholinergies.

#### Kits/Articles of Manufacture

For use in the therapeutic methods of use described 20 herein, kits and articles of manufacture are also described herein. Such kits include a carrier, package, or container that is compartmentalized to receive one or more containers such as vials, tubes, and the like, each of the container(s) comprising one of the separate elements to be used in a method 25 described herein. Suitable containers include, for example, bottles, vials, syringes, and test tubes. In one embodiment, the containers are formed from a variety of materials such as glass or plastic.

The articles of manufacture provided herein contain packaging materials. Packaging materials for use in packaging pharmaceutical products include, e.g., U.S. Pat. No. 5,323, 907. Examples of pharmaceutical packaging materials include, but are not limited to, blister packs, bottles, tubes, bags, containers, bottles, and any packaging material suitable for a selected formulation and intended mode of administration and treatment.

In some embodiments, the compounds or compositions described herein, are presented in a package or dispenser device which may contain one or more unit dosage forms 40 containing the active ingredient. The compound or composition described herein is packaged alone, or packaged with another compound or another ingredient or additive. In some embodiments, the package contains one or more containers filled with one or more of the ingredients of the pharma- 45 ceutical compositions. In some embodiments, the package comprises metal or plastic foil, such as a blister pack. In some embodiments, the package or dispenser device is accompanied by instructions for administration, such as instructions for administering the compounds or composi- 50 tions for treating a neoplastic disease. In some embodiments, the package or dispenser is accompanied with a notice associated with the container in form prescribed by a governmental agency regulating the manufacture, use, or sale of pharmaceuticals, which notice is reflective of approval by 55 the agency of the form of the drug for human or veterinary administration. In some embodiments, such notice, for example, is the labeling approved by the U.S. Food and Drug Administration for prescription drugs, or the approved product insert. In some embodiments, compositions include 60 a compound described herein formulated in a compatible pharmaceutical carrier are prepared, placed in an appropriate container, and labeled for treatment of an indicated condition.

For example, the container(s) include Compound 1, 65 optionally in a composition or in combination with another agent as disclosed herein. Such kits optionally include an

86

identifying description or label or instructions relating to its use in the methods described herein.

A kit typically includes labels listing contents and/or instructions for use, and package inserts with instructions for use. A set of instructions will also typically be included.

In one embodiment, a label is on or associated with the container. In one embodiment, a label is on a container when letters, numbers or other characters forming the label are attached, molded or etched into the container itself; a label is associated with a container when it is present within a receptacle or carrier that also holds the container, e.g., as a package insert. In one embodiment, a label is used to indicate that the contents are to be used for a specific therapeutic application. The label also indicates directions for use of the contents, such as in the methods described herein.

In certain embodiments, the pharmaceutical compositions are presented in a pack or dispenser device which contains one or more unit dosage forms containing a compound provided herein. The pack, for example, contains metal or plastic foil, such as a blister pack. In one embodiment, the pack or dispenser device is accompanied by instructions for administration. In one embodiment, the pack or dispenser is also accompanied with a notice associated with the container in form prescribed by a governmental agency regulating the manufacture, use, or sale of pharmaceuticals, which notice is reflective of approval by the agency of the form of the drug for human or veterinary administration. Such notice, for example, is the labeling approved by the U.S. Food and Drug Administration for prescription drugs, or the approved product insert. In one embodiment, compositions containing a compound provided herein formulated in a compatible pharmaceutical carrier are also prepared, placed in an appropriate container, and labeled for treatment of an indicated condition.

### **EXAMPLES**

The following ingredients, formulations, processes and procedures for practicing the methods disclosed herein correspond to that described above.

Example 1: Wet Granulation Method for the Preparation of High-Load Tablet Formulations of Ibrutinib

A high shear mixer is charged with ibrutinib, intragranular components, such as lactose monohydrate, optionally microcrystalline cellulose, and optionally croscarmellose sodium and hydroxypropylcellulose, in W/W proportions as described in Tables 1A-1F. The ingredients were then mixed and water (or an aqueous binder solution) was added gradually. Once granulated, the wet granules were dried in a fluid bed dryer with an inlet temperature at 60° C. until the loss on drying (LOD) was <1.5%. The dried granulated material was then passed through a comil equipped with a 0.04 inch screen. The extragranular components, such as croscarmellose sodium, sodium lauryl sulfate and colloidal silicon dioxide were then mixed with the granules in a blender for 20 minutes. Magnesium stearate was charged to the blender. The final mixture was blended for another two minutes. The final blend was then discharged and compressed into tablets. The tablets are stored at room temperature until they are used.

Wet granulation tablet formulations W1-W19, W22, W23-W24, and BK01-BK02, BK04, BK06-BK09, BK21A

and BK21B shown below in Tables 1A-1F were prepared as or similarly to that described above.

TABLE 1A

			V	Vet Grai	nulation 7	Tablet F	ormulat	ions						
		/1	W	/2	W	3	w	<u> 4</u>	w	<u> 5</u>	w	<u> 16</u>		v7
	Wt %	mg/ Tab	Wt %	mg/ Tab	Wt %	mg/ Tab	Wt %	mg/ Tab	Wt %	mg/ Tab	Wt %	mg/ Tab	Wt %	mg/ Tab
Ibrutinib	60.0	560.0	60.0	560.0	60.0	560.0	60.0	560.0	60.0	560.0	60.0	560.0	70.0	560.0
MCC (Avicel PH101)	12.5	116.7	12.5	116.7	10.75	100.3	25.0	233.3	_		12.5	116.7	15.0	120.0
Lactose Mono (Fast	12.5	116.6	12.5	116.6	10.75	100.3	_	_	25.0	233.3	_	_	_	_
Flo 316)											12.5	1166		
Mannitol (Pearlitol 100 SD)			_		_	_	_	_	_		12.5	116.6	_	_
Hydroxypropylcellulose (Klucel EXF)	3.0	28.0	3.0	28.0	_	_	3.0	28.0	3.0	28.0	3.0	28.0	3.0	24.0
PVP K30	_	_	_	_	6.5	60.7	_	_		_	_	_	_	
SLS (Kolliphor Fine)		_	3.0	28.0		_	_	_				_	_	_
Subtotal (Intra)	88.0	821.3	91.0	849.3	88.0	821.3	88.0	821.3	88.0	821.3	88.0	821.3	88.0	704.0
SLS (Kolliphor Fine)	6.0	56.0	3.0	28.0	6.0	56.0	6.0	56.0	6.0	56.0	6.0	56.0	6.0	48.0
Croscarmellose Na (Ac-	5.0	46.7	5.0	46.7	5.0	46.7	5.0	46.7	5.0	46.7	5.0	46.7	5.0	40.0
Di-Sol)	5.0	10.7	5.0	10.7	5.0	10.7	2.0	10.7	5.0	10.7	5.0	10.7	5.0	10.0
Silicon Dioxide	0.5	4.7	0.5	4.7	0.5	4.7	0.5	4.7	0.5	4.7	0.5	4.7	0.5	4.0
(Cabosil M5P)														
Magnesium Stearate	0.5	4.6	0.5	4.6	0.5	4.6	0.5	4.6	0.5	4.6	0.5	4.6	0.5	4.0
Total	100	933.3	100	933.3	100	933.3	100	933.3	100	933.3	100	933.3	100%	800.0

TABLE 1B

<b>W</b> 10	W9	
		W11
60.0%	70.0%	70.0%
13.5%	9.0%	8.5%
13.5%	9.0%	8.5%
2.5%	_	3.0%
1.0%		
90.5%	88.0%	90.0%
6.0%	6.0%	6.0%
2.5%	5.0%	3.0%
0.5%	0.5%	0.5%
0.5%	0.5%	0.5%
	0.5%	

TABLE 1C

Wet G	ranulation	Tablet For	mulations	!	
	W12	W13	W14	W15	W16
Ibrutinib	60.0%	60.0%	70.0%	70.0%	70.0%
MCC (Avicel PH101)	12.0%	12.0%	7.0%	7.0%	_
Lactose Mono (Fast Flo 316)	12.0%	12.0%	7.0%	7.0%	14.0%
Croscarmellose Na (Ac-Di-Sol)	5.0%	5.0%	5.0%	5.0%	5.0%
Hydroxypropylcellulose (SSL)	2.0%	_	2.0%	_	_
Povidone (K25)		2.0%		2.0%	2.0%
Subtotal (Intra)			91.0%		
SLS (Kolliphor Fine)	6.0%	6.0%	6.0%	6.0%	6.0%
Croscarmellose Na (Ac-Di-Sol)	2.0%	2.0%	2.0%	2.0%	2.0%

## TABLE 1C-continued

Wet Granulation Tablet Formulations										
	W12	W13	W14	W15	W16					
Silicon Dioxide (Cabosil M5P)	0.5%	0.5%	0.5%	0.5%	0.5%					
Magnesium Stearate	0.5%	0.5%	0.5%	0.5%	0.5%					
Total			100.0%							

## TABLE 1D

		Wet	Granulation	Tablet Form	nulations			
			Blend	_				
Component		W17		W18	W19	W21	W23	W24
Intragranular	_							
Ibrutinib Lactose (Fast Flo 316)		70.00% 14.00%		70.00% 14.00%	70.00% 14.00%	70% 13.00%	70% 13.00%	70% 14.00%
PVP K25 Croscarmellose Na (Ac-Di-Sol)		2.00% 5.00%		2.00% 5.00%	2.00% 5.00%	3.00% 5.00%	3.00% 5.00%	2.00% 5.00%
SLS (Kolliphor Fine) Poloxamer		1.00%		0.00%	0.00%	1.00%	1.00%	1.00%
Tween 80				0.50%	1.0070			
Subtotal (Intra) Water		92.00% 35%		91.50% 35%	92.00% 35%	92.00% 35%	92.00% 40%	92.00% 40%
Extragranular	17A	17B	17C	_				
Lactose (Fast Flo 316)	5.00%	0.00%	0.00%	5.50%	5.00%	5.00%	5.00%	5.00%
Citric Acid		5.00%						
Poloxamer Croscarmellose Na (Ac-Di-Sol)	2.00%	2.00%	5.00% 2.00%	2.00%	2.00%	2.00%	2.00%	2.00%
Silicon Dioxide (Cabosil M5P)	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%
Magnesium Stearate	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%
Total	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%

TABLE 1E

			Wet Granu	lation Tablet	Formulatio:	ns			
Component	ВК	01	BK02	BK04	BK06	BK07	BK08	BK09	BK10
Intragranular	_								
Ibrutinib Lactose (Fast Flo 316)		00% 00%	70.00% 14.00%	70.00% 14.00%	70.00% 13.00%	70% 14.00%	70% 6.00%	70% 5.00%	70% 16%
Povidone (PVP K25)	2.	00%	2.00%	2.00%	3.00%	2.00%	2.00%	2.00%	2.0%
Cropovidone Sodium Starch	5.	00%	10.00%	0	5.00%	5.00%	15.0%	15.0%	10.0%
Glycolate SLS (Kolliphor Fine)	0	00%	0 1.00%	5.00% 0.00%	0 1.0%	0 3.00%	0 1.00%	0 3.00%	0 1.0%
Subtotal (Intra) Extragranular	92. BK01A	00% BK01B	97.00%	92.0%	92.00%	94.00%	94.00%	95.00%	99.0%
Lactose (Fast Flo 316)	5.00%	2.00%	2.00%	5.0%	5.0%	3.00%	5.00%	4.00%	0
Crospovidone	2.00%	5.00%	0	0	2.00%	2.00%	0	0	0
Sodium Starch Glycolate	0	0	0	2.00%	0	0	0	0	0
Silicon Dioxide (Cabosil M5P)	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%
Magnesium Stearate	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%	0.50%
Total	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.0%

Ingredients

Ibrutinib (Intra)

Lactose Monohydrate (Intra)

Binder (PVP K25) (Intra)

SLS (% Intra/% Extra)

Mg Stearate (Extra)

Microcrystalline Cellulose (Extra)

Colloidal Silicon Dioxide (Extra)

Croscarmellose Sodium (% intra/Extra)

BK21A

70%

14%

5%

2%

1% (1/0)

7% (5/2)

0.5%

BK21B

70%

14%

4% (1/3)

7% (5/2)

0.5%

2%

92
TABLE 3-continued

	Components of Imme	rdiate Release Tablet Formulation
5	Ingredient	Range
, <b>-</b>	Microcrystalline cellulose Colloidal Silicon Dioxide Magnesium stearate Total	0 to 10% 0 to 1% 0.25% to 2.5% Tablet weight range: 622 mg to 700 mg
10		

Example 4: Preparation of Capsule Formulations of Ibrutinib

Example 2: Dry Granulation Method for the Preparation of Ibrutinib Tablet Formulations

Ibrutinib, microcrystalline cellulose, croscarmellose sodium, sodium lauryl sulfate and optionally magnesium stearate were passed through a 1000 microns sieve. The 20 mixture was then blended for 10 minutes. The pre-blending was charged to a roller compactor and compacted at 0.6 kN/cm. The resulting ribbon was passed through an oscillating mill equipped with a 0.8 mm screen. The milled granules were then combined with the extragranular components: microcrystalline cellulose, croscarmellose sodium, sodium lauryl sulfate and optionally magnesium stearate and blended for 10 minutes. The blend was then compressed into tablets using a single station manual press.

Dry granulation tablet formulations D1 and D5 shown <sup>30</sup> below in Table 2 were prepared as described above.

TABLE 2

	<u>-</u>								
Dry Granulation Tablet Formulations									
		D1							
Component	Wt %	mg/Tab	Wt %	mg/Tab					
Ibrutinib	50.00	560.0	42.42	560.0	40				
MCC (Avicel PH101)	30.00	336.0	45.88	605.7	40				
Croscarmellose Na (Ac-Di-Sol)	4.00	44.8	3.97	52.4					
SLS (Kolliphor Fine)	3.40	38.1	2.85	37.6					
Magnesium Stearate			0.24	3.2					
Subtotal (Intra)	87.40	978.9	95.36	1258.9					
MCC (Avicel PH200)	7.50	84.0	_	_	45				
SLS (Kolliphor Fine)	1.60	17.9	1.39	18.3					
Croscarmellose Na (Ac-Di-Sol)	3.00	33.6	3.00	39.6					
Magnesium Stearate	0.50	5.6	0.24	3.2	-				
Total	100.00	1120.0	100.00	1320.1	50				

Example 3: Preparation of Immediate Release High-Load Tablet Formulations of Ibrutinib

Immediate release tablet formulations are prepared using the components shown in Table 3 following the procedure from Example 1.

TABLE 3

60

Components of Immediate Release Tablet Formulation						
Ingredient	Range					
Ibrutinib	80 to 90%					
Lactose	0 to 10%					
Croscarmellose sodium	1 to 10%					

The capsule Formulation A manufacturing process includes the following steps: weigh the indicated amount of the components, mix together and add into an appropriate size capsule, and close capsule. The capsules are stored at room temperature until they are used.

Example 5: In Vivo Evaluation of High-Load, Wet and Dry Granulation Tablet Formulations of Ibrutinib and Ibrutinib Capsule Formulation

The pharmacokinetics of ibrutinib in capsule (Formulation A) versus different wet granulation (Formulations B, C and D which correspond to Formulations W8, W10, and W11 in Example 1, respectively) and dry granulation (Formulations E and F which correspond to Formulations D1 and D5 in Example 2, respectively) tablet formulations was studied in fasted male beagle dogs following single oral administration of 140 mg ibrutinib formulations administered in a Latin square crossover design. The wet granulation (Formulations E and F) tablets were studied in two parallel groups with capsule as internal control in each of the group. The tablet formulations composition and drug load are presented in Table 4.

TABLE 4

<u>Ibrutinib Ta</u>	blet Form	ulations -	Drug Loa Formul		nposition	
	A	В	C Proce	D	E	F
Component	— w/w %	Wet w/w %	Wet w/w %	Wet w/w %	Dry w/w %	Dry w/w %
Ibrutinib Lactose Mono- hydrate NF	42.0 0	60.0 14	60.0 13.5	70.0 8.5	50.0 0	<b>42.</b> 0
Microcrystalline cellulose NF	46.5	14	13.5	8.5	37.5	46.5
Hydroxypropyl Cellulose NF	0	0	1.0	0	0	0
Croscarmellose sodium NF	7.0	5.0	5.0	6.0	7.0	7.0
Sodium lauryl sulfate NF	4.0	6.0	6.0	6.0	5.0	4.0
Colloidal Silicon Dioxide NF	0	0.5	0.5	0.5	0	0
Magnesium stearate NF	0.5	0.5	0.5	0.5	0.5	0.5
Tablet Weight	333.3	233.3	233.3	200.0	280.0	333.3

FIG. 1 and FIG. 2 shows mean plasma concentration-time profiles of ibrutinib following single oral dose administra-

formulations, respectively to fasted beagle dogs (Dose=140 mg). In general, all the wet granulation formulation tablets

(B, C and D) tested showed comparable concentrations to

the capsule formulation. Specifically, wet granulation tablet formulations B, C and D showed average % Frel values ranging from 72 to 110% (Table 5). However, dry granulation tablet formulations, E and F showed lower concentrations compared to capsule with average % F<sub>red</sub> values ranging from 43 to 52% (Table 6). Furthermore, reduced variability in ibrutinib exposure was observed with wet granulation formulations when compared to capsule formulation (Tables 5 and 7). In some embodiments, the tablet formulation provides unexpected low variability in ibrutinib exposure in terms of %  $\overline{\text{CV}}$  for both  $C_{max}$  and AUC when 94

TABLE 5-continued

Mean (% CV) Ibrutinib Plasma PK Parameters Following Single Dose Administration of Three Different Wet Tablet Ibrutinib Formulations to Fasted Beagle Dogs (n = 7)

	Formulation	Dose (mg)	C <sub>max</sub> (ng/mL)	$\begin{array}{c} T_{max} \\ (h) \end{array}$	T <sub>1/2</sub> (h)	AUC (ng * h/mL)	$\mathbf{F}_{rel}\left(\%\right)$
0	B (Wet W8)	140	20.7 (54.1)	1.64	ND	63.2 (37.7)	96 (63.1)
	C (Wet W10)	140	22.6 (86.1)	1.46	1.10 <sup>a</sup>	72.6 (66.1)	72 (53.2)
	D (Wet W11)	140	23.0 (86.9)	2.00	6.50 <sup>b</sup> (36.5)	93.3 (92.8)	110 (83.3)

 $F_{ref.}$  for each dog, which has received both capsule and tablet formulations in a cross-over fashion (AUC<sub>Formulation B, C, D, E or F/AUC<sub>Formulation A</sub>) \* 100;</sub>

N/A: not applicable;

 $^{b}$ n = 3;

ND: not determined

TABLE 5

without food, see, e.g., Table 7.

administered under fasted and fed conditions, see, e.g., Formulation BK21B as compared to the capsule formulation

A and tablet formulations BK02 and BK21A (Table 7). In

some embodiments, the tablet formulation provides unex-

pected absence of food effect in  $C_{max}$  and  $T_{max}$  when administered under fed conditions compared to fasted con-

ditions, which is contemplated to lead to more predictable therapeutic efficacy and side effects when taking with or

Mean (% CV) Ibrutinib Plasma PK Parameters Following Single Dose Administration of Three Different Wet Tablet Ibrutinib Formulations to Fasted Beagle Dogs (n = 7)

Formulation	Dose (mg)	C <sub>max</sub> (ng/mL)		T <sub>1/2</sub> (h)	AUC (ng * h/mL)	$\mathbf{F}_{rel}  (\%)$
A (Capsule)	140	32.1 (93.7)	2.36	ND	146 (106)	N/A

TABLE 6

Mean (% CV) Ibrutinib Plasma PK Parameters Following Single Dose Administration of Two Different Dry Tablet Ibrutinib Formulations to Fasted Beagle Dogs (n = 8)

Formulation	Dose (mg)	$\begin{array}{c} {\rm C}_{max} \\ ({\rm ng/mL}) \end{array}$	$\begin{array}{c} \mathbf{T}_{max} \\ (\mathbf{h}) \end{array}$	T <sub>1/2</sub> (h)	AUC (ng * h/mL)	$\mathbf{F}_{rel}\left(\% ight)$
A (Capsule) E (Dry D1) F (Dry D5)	140 140 140	29.9 (90.2) 10.2 (78.9) 10.1 (49.5)	2.72	ND ND ND	161 (105) 45.1 (43.7) 59.9 (77.3)	N/A 42.6 (64.7) 51.6 (66.5)

 $F_{ref}$ : for each dog, which has received both capsule and tablet formulations in a cross-over fashion (AUC Formulation B, C, D, E or FAUC Formulation A) \* 100; N/A: not applicable;

ND: not determined

TABLE 7

Comparison of Capsule and Three Tablet Formulations (140 mg dose) in Dogs under Fasted and Fed States

		$C_{max}$	$T_{max}$	T <sub>1/2</sub>	$\mathrm{AUC}^a$	tablet/	rel capsule	Fed/F	rel fasted 6)
Formu	ılation	(ng/mL)	(h)	(h)	(ng * h/mL)	$C_{max}$	AUC	$C_{max}$	AUC
A (Capsule)	$\mathrm{Fed}^{c,d}$	251 (78.6) <sup>e</sup>	3.20	1.74	892 (70.7)	NA	NA	89.7	117
(	Fasted <sup>f,g</sup>	419 (57.1)	1.63	2.23	1010 (52.7)	NA	NA		
BK02	$\mathrm{Fed}^h$	352 (54.3)	2.00	1.92	991 (42.2)	236	146	84.1	107
	Fasted	494 (54.0)	1.19	2.20	1000 (52.9)	150	112		
BK21A	Fed	287 (75.5)	2.93	1.63	878 (60.5)	172	119	92.1	99.2
	Fasted	370 (73.5)	1.47	2.47	1000 (64.6)	111	108		
BK21B	Fed	367 (51.1)	1.80	1.73	1014 (37.5)	227	146	104	136
	Fasted	363 (45.6)	1.44	1.85	840 (36.3)	152	134		

<sup>&</sup>lt;sup>a</sup>AUC<sub>last</sub>

 $<sup>^{</sup>b}$ calculated for each dog, which has received both capsule and tablet formulations in a cross-over fashion <sup>c</sup>fed dogs were administered with liquid concentrated diet 15 minutes prior to dosing

 $<sup>^</sup>e$ % coefficient of variation (CV), values are presented as mean (% CV)

 $f_{
m fasted}$  dogs were administered 12 mg/kg sub-cutaneous pentagastrin 45 minutes prior to dosing

 $g_{n} = 16$ 

 $<sup>^{</sup>h}$ n = 14

Example 6: A Single-Dose, Open-Label, Randomized, Crossover Study to Assess the Pharmacokinetics of Ibrutinib Tablet Formulations in Healthy Adult Subjects Compared to the Ibrutinib Capsule Formulation a

This is a single-center, open-label, randomized, crossover, single-dose study in healthy adults. After providing written informed consent, subjects were screened within 21 days (Day -21 to -2).

Main Criteria for Inclusion:

 $AUC_{\infty}$ : Area under the concentration-time curve from time 0 to infinite time

t<sub>1/2</sub>: Apparent elimination half-life associated with the terminal slope of the semilogarithmic drug concentration-time curve

TABLE 8

Pharmacokinetics Parameters and Results								
Treatment	N		C <sub>max</sub> (ng/mL)	$\begin{array}{c} T_{max} \\ (h) \end{array}$	$\begin{array}{c} \mathrm{AUC}_{last} \\ (\mathrm{ng}  *  \mathrm{h/mL}) \end{array}$	$\begin{array}{c} \mathrm{AUC}_{inf} \\ (\mathrm{ng} \ * \ \mathrm{h/mL}) \end{array}$	T <sub>1/2</sub> (h)	
Ibrutinib capsule Formulation A (4 × 140 mg/ capsule) Ibrutinib	32	Mean SD Range % CV Mean	48.6 36.0 7.50-184 74.1 47.7	1.64 1.09 0.5-4 66.2	379 248 118-1100 62.4 413	465 <sup>a</sup> 248 206-1120 53.4 472 <sup>b</sup>	9.5 <sup>a</sup> 3.5 5.9-20.0 36.9 8.3 <sup>b</sup>	
Tablet Formulation BK21A coated with a cosmetic film coating agent - Opadry II white (560 mg/ tablet)	22	SD Range % CV	43.7 7.50-181 91.6	4.88 1-24 174	227 135-1040 55.0	246 155-1060 52.0	1.9 5.9-13.1 22.8	
Ibrutinib Tablet Formulation BK21B coated with a cosmetic film coating agent - Opadry II white (560 mg/ tablet)	21	Mean SD Range % CV	35.5 21.9 7.90-96.0 61.6	1.90 1.28 1-6 67.2	355 135 74.2-692 38.1	411° 130 182-696 31.7	7.8° 2.0 5.3-13.2 26.1	

 $a_n = 22;$ 

Healthy men and women between 18 and 55 years of age, inclusive; body mass index (BMI) between 18 and 30 kg/m², inclusive, and a body weight of not less than 50 kg. Women must be post-menopausal or surgically sterile.

Eligible subjects received a single oral dose of ibrutinib 560 mg (in either capsule formulation A comprising 140 mg ibrutinib per capsule or a tablet formulation comprising 560 mg ibrutinib per tablet) with 240 mL of noncarbonated water 50 on Day 1 of each treatment period after fasting at least 10 hours before each dose. Water was allowed ad libitum beginning 2 hours after each dose, and lunch was provided beginning 4 hours after each dose.

Blood samples for pharmacokinetic (PK) analysis of 55 ibrutinib were collected before dosing and over 48 hours after dosing in each treatment period.

Total duration of the study was approximately 70 days (21-day screening period, 4×3-day treatment periods with 7-day washouts between periods, and a 7-day follow-up 60 phase).

PK parameters including the following were calculated and the results are in Table 8:

C<sub>max</sub>: Maximum observed concentration

T<sub>max</sub>: Time to reach the maximum observed concentration 65
 AUC<sub>tast</sub>: Area under the concentration-time curve from time 0 to last time point

In some embodiments, the high load tablet formulations possess both pharmaceutically acceptable properties and desired PK properties, such as a high  $C_{max}$ , similar to that of a capsule formulation (e.g., BK21A).

Example 7: Safety and Tolerability Study of Compound 1 in Chronic Lymphocytic Leukemia

Purpose: The purpose of this study is to establish the safety and optimal dose of orally administered Compound 1 (420 mg/day) high-load tablets in patients with B-cell chronic lymphocytic leukemia/small lymphocytic lymphoma/diffuse well-differentiated lymphocytic lymphoma.

Primary Outcome Measures: Safety and tolerability of Compound 1 (frequency, severity, and relatedness of adverse events).

Secondary Outcome Measures: Pharmacokinetic/Pharmacodynamic assessments. Tumor response—overall response rate as defined by recent guidelines on CLL and SLL (B cell lymphoma) and duration of response.

Eligibility: 18 Years and older; both genders are eligible. Inclusion Criteria: 1. For treatment-naive group only: Men and women ≥65 years of age with confirmed diagnosis of CLL/SLL, who require treatment per NCI or International

 $<sup>^{</sup>b}$ n = 16;

 $c_{n} = 13$ 

97

Working Group guidelines 11-14. 2. For relapsed/refractory group only: Men and women ≥18 years of age with a confirmed diagnosis of relapsed/refractory CLL/SLL unresponsive to therapy (ie, failed ≥2 previous treatments for CLL/SLL and at least 1 regimen had to have had a purine 5 analog [eg, fludarabine] for subjects with CLL). 3. Body weight  $\ge 40$  kg. 4. ECOG performance status of  $\le 2$ . 5. Agreement to use contraception during the study and for 30 days after the last dose of study drug if sexually active and able to bear children. 6. Willing and able to participate in all 10 required evaluations and procedures in this study protocol including swallowing tablets without difficulty. 7. Ability to understand the purpose and risks of the study and provide signed and dated informed consent and authorization to use protected health information (in accordance with national 15 and local subject privacy regulations).

Exclusion Criteria: 1. A life-threatening illness, medical condition or organ system dysfunction which, in the investigator's opinion, could compromise the subject's safety, interfere with the absorption or metabolism of Compound 1 20 PO, or put the study outcomes at undue risk. 2. Any immunotherapy, chemotherapy, radiotherapy, or experimental therapy within 4 weeks before first dose of study drug (corticosteroids for disease-related symptoms allowed but require 1-week washout before study drug administration). 25 3. Central nervous system (CNS) involvement by lymphoma. 4. Major surgery within 4 weeks before first dose of study drug. 5. Creatinine >1.5×institutional upper limit of normal (ULN); total bilirubin >1.5×ULN (unless due to Gilbert's disease); and aspartate aminotransferase (AST) or 30 alanine aminotransferase (ALT) >2.5×ULN unless disease related. 6. Concomitant use of medicines known to cause QT prolongation or torsades de pointes. 7. Significant screening electrocardiogram (ECG) abnormalities including left bundle branch block, 2nd degree AV block type II, 3rd 35 degree block, bradycardia, and QTc >470 msec. 8. Lactating or pregnant.

## Example 8: Safety and Efficacy of Compound 1 in Subjects with Relapsed/Refractory Mantle Cell Lymphoma (MCL)

The primary objective of this trial is to evaluate the efficacy of Compound 1 in relapsed/refractory subjects with Mantle Cell Lymphoma (MCL). The secondary objective is 45 to evaluate the safety of a fixed daily dosing regimen of Compound 1 (560 mg/day in the form of tablets) in this population.

Primary Outcome Measures: To measure the number of participants with a response to Compound 1.

Secondary Outcome Measures: To measure the number of participants with adverse events as a measure of safety and tolerability. To measure pharmacokinetics to assist in determining how the body responds to the study drug. Patient reported outcomes (to measure the number of participants 55 420 mg tablet) orally daily and will be continued daily. reported outcomes in determining the health related quality

Eligibility: 18 Years and older; both genders are eligible. Inclusion Criteria: Men and women ≥18 years of age. ECOG performance status of ≤2. Pathologically confirmed 60 MCL, with documentation of either overexpression of cyclin D1 or t(11;14), and measurable disease on cross sectional imaging that is ≥2 cm in the longest diameter and measurable in 2 perpendicular dimensions. Documented failure to achieve at least partial response (PR) with, or documented 65 disease progression disease after, the most recent treatment regimen. At least 1, but no more than 5, prior treatment

98

regimens for MCL (Note: Subjects having received ≥2 cycles of prior treatment with bortezomib, either as a single agent or as part of a combination therapy regimen, will be considered to be bortezomib-exposed.). Willing and able to participate in all required evaluations and procedures in this study protocol including swallowing tablets without difficulty. Ability to understand the purpose and risks of the study and provide signed and dated informed consent and authorization to use protected health information (in accordance with national and local subject privacy regulations).

Major exclusion criteria: Prior chemotherapy within 3 weeks, nitrosoureas within 6 weeks, therapeutic anticancer antibodies within 4 weeks, radio- or toxin-immunoconjugates within 10 weeks, radiation therapy within 3 weeks, or major surgery within 2 weeks of first dose of study drug. Any life-threatening illness, medical condition or organ system dysfunction which, in the investigator's opinion, could compromise the subject's safety, interfere with the absorption or metabolism of Compound 1 high-load tablets, or put the study outcomes at undue risk. Clinically significant cardiovascular disease such as uncontrolled or symptomatic arrhythmias, congestive heart failure, or myocardial infarction within 6 months of screening, or any Class 3 or 4 cardiac disease as defined by the New York Heart Association Functional Classification. Malabsorption syndrome, disease significantly affecting gastrointestinal function, or resection of the stomach or small bowel or ulcerative colitis, symptomatic inflammatory bowel disease, or partial or complete bowel obstruction. Any of the following laboratory abnormalities: 1. Absolute neutrophil count (ANC)<750 cells/mm3 (0.75×109/L) unless there is documented bone marrow involvement. 2. Platelet count <50,000 cells/mm3 (50×109/L) independent of transfusion support unless there is documented bone marrow involvement. 3. Serum aspartate transaminase (AST/SGOT) or alanine transaminase (ALT/SGPT) ≥3.0×upper limit of normal (ULN). 4. Creatinine  $>2.0\times ULN$ .

Example 9: Phase 2 Study of the Combination of Compound 1 and Rituximab in High-Risk Chronic Lymphocytic Leukemia and Small Lymphocytic Lymphoma Patients

Purpose: The goal of this clinical research study is to learn if Compound 1 combined with rituximab can help to control chronic lymphocytic leukemia (CLL) and small lymphocytic lymphoma (SLL). The safety of this combination will 50 also be studied.

Rituximab (375 mg/m<sup>2</sup>) will be given intravenously (IV) on Day 1, Day 8, Day 15, and Day 22, then continued once every 4 weeks only on Days 1 during cycles 2-6. Compound 1 will be started on Day 2 of cycle 1 at a dose of 420 mg (one

Primary Outcome Measures: Progression free survival (PFS) [Time Frame: 3 months]—progression free survival defined as the time interval from treatment to progressive disease or death, whichever happens earlier. Patients in complete remission (CR), partial remission (PR) or stable disease (SD) are all counted as progression-free. Survival or times to progression functions estimated using the Kaplan-Meier method.

Secondary Outcome Measures: Toxicity [Time Frame: 3 months]—toxicity reported by type, frequency and severity. Worst toxicity grades per patient tabulated for selected adverse events and laboratory measurements. Toxicity

(grade 3 or 4) monitored based on the Bayesian model (beta-binomial) by assuming a priori probability of toxicity following beta(1,1).

Eligibility: 18 Years and older; both genders are eligible. Inclusion Criteria: 1. Patients must have a diagnosis of 5 high-risk CLL/SLL and be previously treated with up to 3 lines of prior therapy. High-risk CLL and high-risk SLL is defined by the presence of a 17p deletion or 11q deletion or TP53 mutation. Any CLL and SLL patient who has a short remission duration of less than 3 years after prior first-line 10 chemo-immunotherapy, such as the FCR regimen, also fulfills criteria of high-risk CLL/SLL, regardless of the presence or absence of cytogenetic abnormalities. 2. CLL and SLL patients with 17p deletion or TP53 mutation will not be required to have received any prior therapy, given the poor outcome of CLL/SLL patients to standard frontline chemo-immunotherapy, such patients will be eligible if they are untreated or if they have received up to 3 lines of prior therapy. 3. Patients must have an indication for treatment by 2008 IWCLL Criteria. 4. Patients age >18 years at the time 20 of signing informed consent. Understand and voluntarily sign an informed consent. Be able to comply with study procedures and follow-up examinations. 5. ECOG/WHO performance status of 0-1. 6. Patients of childbearing potential must be willing to practice highly effective birth control 25 (e.g., condoms, implants, injectables, combined oral contraceptives, some intrauterine devices [IUDs], sexual abstinence, or sterilized partner) during the study and for 30 days after the last dose of study drug. Women of childbearing potential include any female who has experienced menarche 30 and who has not undergone successful surgical sterilization (hysterectomy, bilateral tubal ligation, or bilateral oophorectomy) or is not postmenopausal. Post menopause is defined as follows: Amenorrhea >/=12 consecutive months without another cause and a documented serum follicle stimulating 35 hormone (FSH) level >35 mIU/mL; a male of childbearing potential is any male that has not been surgically sterilized. 7. Adequate renal and hepatic function as indicated by all of the following: Total bilirubin </=1.5xinstitutional Upper Limit of Normal (ULN) except for patients with bilirubin 40 elevation due to Gilbert's disease who will be allowed to participate; an ALT </=2.5×ULN; and an estimated creatinine clearance (CrCl) of >30 mL/min, as calculated by the Cockroft-Gault equation unless disease related. 8. Free of prior malignancies for 3 years with exception of currently 45 treated basal cell, squamous cell carcinoma of the skin, or carcinoma in situ of the cervix or breast. 9. A urine pregnancy test (within 7 days of Day 1) is required for women with childbearing potential.

Exclusion Criteria: 1. Pregnant or breast-feeding females. 50 or more pharmaceutically acceptable excipients, 2. Treatment including chemotherapy, chemo-immunotherapy, monoclonal antibody therapy, radiotherapy, highdose corticosteroid therapy (more than 60 mg Prednisone or equivalent daily), or immunotherapy within 21 days prior to enrollment or concurrent with this trial. 3. Investigational 55 agent received within 30 days prior to the first dose of study drug or have previously taken Compound 1. If received any investigational agent prior to this time point, drug-related toxicities must have recovered to Grade 1 or less prior to first dose of study drug. 4. Systemic fungal, bacterial, viral, or 60 other infection not controlled (defined as exhibiting ongoing signs/symptoms related to the infection and without improvement, despite appropriate antibiotics or other treatment). 5. Patients with uncontrolled Autoimmune Hemolytic Anemia (AIHA) or autoimmune thrombocytopenia (ITP). 6. 65 Patients with severe hematopoietic insufficiency, as defined by an absolute neutrophil count of less than 500/micro-L

100

and/or a platelet count of less than 30,000/micro-L at time of screening for this protocol. 7. Any other severe concurrent disease, or have a history of serious organ dysfunction or disease involving the heart, kidney, liver or other organ system that may place the patient at undue risk to undergo therapy with Compound 1 and rituximab. 8. Significant cardiovascular disease such as uncontrolled or symptomatic arrhythmias, congestive heart failure, or myocardial infarction within 6 months of screening, or any Class 3 or 4 cardiac disease as defined by the New York Heart Association Functional Classification. 9. Significant screening ECG abnormalities including left bundle branch block, 2nd degree AV block type II, 3rd degree block, bradycardia, and QTc >470 msec. 10. Any serious medical condition, laboratory abnormality, or psychiatric illness that places the subject at unacceptable risk if he/she were to participate in the study. 11. History of stroke or cerebral hemorrhage within 6 months. 12. Evidence of bleeding diathesis or coagulopathy. 13. Major surgical procedure, open biopsy, or significant traumatic injury within 28 days prior to Day 1, anticipation of need for major surgical procedure during the course of the study. 14. Minor surgical procedures, fine needle aspirations or core biopsies within 7 days prior to Day 1. Bone marrow aspiration and/or biopsy are allowed. 15. Serious, non-healing wound, ulcer, or bone fracture. 16. Treatment with Coumadin. Patients who recently received Coumadin must be off Coumadin for at least 7 days prior to start of the study. 17. Any chemotherapy (e.g., bendamustine, cyclophosphamide, pentostatin, or fludarabine), immunotherapy (e.g., alemtuzumab, or ofatumumab), bone marrow transplant, experimental therapy, or radiotherapy is prohibited during therapy on this study. 18. Use of medications known to prolong QTc interval or that may be associated with Torsades de Pointes (refer to Appendix F) are prohibited within 7 days of starting study drug and during study-drug treatment.

The examples and embodiments described herein are illustrative and various modifications or changes suggested to persons skilled in the art are to be included within this disclosure. As will be appreciated by those skilled in the art, the specific components listed in the above examples may be replaced with other functionally equivalent components, e.g., diluents, binders, lubricants, fillers, and the like.

What is claimed is:

1. A solid tablet formulation comprising ibrutinib and one

wherein ibrutinib is a compound with the structure of Compound 1,

Compound 1

wherein the ibrutinib is in micronized form; and wherein ibrutinib is in an amount of about 140 mg, about 280 mg, about 420 mg, or about 560 mg in the tablet.

- 2. The solid tablet formulation of claim 1, wherein the particle size of micronized ibrutinib is about or less than 30 microns.
- 3. The solid tablet formulation of claim 1, wherein the particle size of micronized ibrutinib is about or less than 10  $_{\rm 30}$  microns.
- **4**. The solid tablet formulation of claim **1**, wherein the particle size of micronized ibrutinib is less than about 1 micron.
- **5**. The solid tablet formulation of claim **1** further comprising one or more diluents.
- 6. The solid tablet formulation of claim 5, wherein the one or more diluents are selected from the group consisting of lactose, sucrose, dextrose, dextrates, maltodextrin, mannitol, xylitol, sorbitol, cyclodextrins, calcium phosphate, calcium sulfate, starches, modified starches, cellulose, microcrystalline cellulose, microcellulose, and talc.
- 7. The solid tablet formulation of claim 5, wherein the one or more diluents are lactose, cellulose, microcrystalline cellulose, or a combination thereof.
- **8**. The solid tablet formulation of claim **5** further comprising one or more lubricants.
- **9**. The solid tablet formulation of claim **8**, wherein the one or more lubricants are a stearate, a polyethylene glycol (PEG), or a wax.
- 10. The solid tablet formulation of claim 8 further comprising one or more disintegrating agents.
- 11. The solid tablet formulation of claim 10, wherein the one or more disintegrating agents are selected from the group consisting of natural starch, a pregelatinized starch, a 55 sodium starch, methylcrystalline cellulose, methylcellulose, croscarmellose, sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymethylcellulose, cross-linked croscarmellose, cross-linked starch, cross-linked polymer, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum.
- 12. The solid tablet formulation of claim 10, wherein the one or more disintegrating agents are selected from the group consisting of natural starch, a pregelatinized starch, a sodium starch, methylcrystalline cellulose, methylcellulose, 65 croscarmellose, croscarmellose sodium, cross-linked sodium carboxymethylcellulose, cross-linked carboxymeth-

102

ylcellulose, cross-linked croscarmellose, sodium starch glycolate, crospovidone, cross-linked polyvinylpyrrolidone, sodium alginate, a clay, and a gum.

- 13. The solid tablet formulation of claim 8 further comprising one or more binders.
- **14**. The solid tablet formulation of claim **13**, wherein the one or more binders are polyvinylpyrrolidone or hydroxypropyl cellulose.
- **15**. The solid tablet formulation of claim 1 comprising: a. at least 50% w/w of ibrutinib;
- b. one or more diluents; and
- c. one or more lubricants.
- **16**. The solid tablet formulation of claim **1** comprising: a. about 50% w/w to about 90% w/w of ibrutinib;
- b. about 5% w/w to about 20% w/w of one or more diluents; and
- c. about 0.1% w/w to about 1.5% w/w of one or more lubricants.
- 17. The solid tablet formulation of claim 1, wherein the excipients comprise intragranular and extragranular excipients; and the intragranular excipients comprise one or more diluents; and the extragranular excipients comprise one or more lubricants.
- 18. The solid tablet formulation of claim 1, wherein the excipients comprise intragranular and extragranular excipients; and the intragranular excipients comprise lactose and microcrystalline cellulose; and the extragranular excipients comprise magnesium stearate.
- 19. The solid tablet formulation of claim 1, wherein the excipients comprise lactose in an amount from about 5% w/w to about 20% w/w, about 10% w/w to about 20% w/w, about 8% w/w to about 15% w/w, about 8.5% w/w to about 14% w/w, or about 12% w/w to about 15% w/w.
- **20**. The solid tablet formulation of claim **1**, wherein the excipients comprise microcrystalline cellulose in an amount from about 5% w/w to about 20% w/w, about 8% w/w to about 20% w/w, about 1% w/w to about 10% w/w, or about 2% w/w to about 5% w/w.
- 21. The solid tablet formulation of claim 1, wherein the excipients comprise magnesium stearate in an amount from about 0.1% w/w to about 1.5% w/w, about 0.4% w/w to about 0.8% w/w, or about 0.5% w/w to about 0.6% w/w.
- 22. The solid tablet formulation of claim 1, wherein the excipients comprise intragranular and extragranular excipients; and the intragranular excipients comprise lactose, microcrystalline cellulose, croscarmellose sodium, and hydroxypropyl cellulose; and the extragranular excipients comprise croscarmellose sodium and magnesium stearate.
- 23. The solid tablet formulation of claim 1, wherein the excipients comprise croscarmellose sodium in an amount from about 0 to about 10% w/w, about 0 to about 5% w/w, about 1% w/w to about 10% w/w, about 2% w/w to about 5% w/w, about 2% w/w to about 4% w/w, about 3% w/w to about 7% w/w, or about 1% w/w to about 3% w/w.
- **24**. The solid tablet formulation of claim 1, wherein the excipients comprise hydroxypropyl cellulose in an amount from about 0 to about 2% w/w, about 0.1% w/w to about 1.1% w/w, or about 0.1% w/w to about 1% w/w.
- 25. The solid tablet formulation of claim 1, wherein the excipients comprise intragranular and extragranular excipients; and the intragranular excipients comprise lactose, polyvinylpyrrolidone, and croscarmellose sodium; and the extragranular excipients comprise croscarmellose sodium, microcrystalline cellulose, and magnesium stearate.

- 26. The solid tablet formulation of claim 1, wherein the excipients comprise polyvinylpyrrolidone in an amount from about 0% w/w to about 5% w/w, or about 1% w/w to about 3% w/w.
- **27**. The solid tablet formulation of claim **1**, wherein the 5 high-load solid tablet formulation is used for one tablet once a day dosing.
- a day dosing.

  28. The solid tablet formulation of claim 1, wherein the formulation comprises about 60% w/w to about 80% w/w of ibrutinib.

\* \* \* \*

# UNITED STATES PATENT AND TRADEMARK OFFICE CERTIFICATE OF CORRECTION

PATENT NO. : 10,010,507 B1 Page 1 of 1

APPLICATION NO. : 15/909779 DATED : July 3, 2018

INVENTOR(S) : Ching W. Chong et al.

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

In the Claims

Column 102,

Claim 23, Line 54: "from about 0 to about 10% w/w, about 0 to about 5% w/w" --from about 0% w/w to about 10% w/w, about 0% w/w to about

5% w/w--

Claim 24, Line 60: "from about 0 to about 2% w/w"

Should read: --from about 0% w/w to about 2% w/w--

Column 103,

Claim 27, Lines 5-6: "the high-load solid tablet formulation"

Should read: --the formulation--

Signed and Sealed this Ninth Day of October, 2018

Andrei Iancu

Director of the United States Patent and Trademark Office